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Gene duplication and genetic exchange drive the evolution of S-RNase-based self-incompatibility in *Petunia*

Kubo, Ken-ichi ; Paape, Timothy ; Hatakeyama, Masaomi ; Entani, Tetsuyuki ; Takara, Akie ; Kajihara, Kie ; Tsukahara, Mai ; Shimizu-Inatsugi, Rie ; Shimizu, Kentaro K ; Takayama, Seiji

Abstract: Self-incompatibility (SI) systems in flowering plants distinguish self- and non-self pollen to prevent inbreeding. While other SI systems rely on the self-recognition between specific male- and female-determinants, the Solanaceae family has a non- self recognition system resulting in the detoxification of female-determinants of S-ribonucleases (S-RNases), expressed in pistils, by multiple male-determinants of S-locus F-box proteins (SLFs), expressed in pollen. It is not known how many SLF components of this non-self recognition system there are in Solanaceae species, or how they evolved. We identified 16–20 SLFs in each S-haplotype in SI *Petunia*, from a total of 168 SLF sequences using large-scale next-generation sequencing and genomic polymerase chain reaction (PCR) techniques. We predicted the target S-RNases of SLFs by assuming that a particular S-allele must not have a conserved SLF that recognizes its own S-RNase, and validated these predictions by transformation experiments. A simple mathematical model confirmed that 16–20 SLF sequences would be adequate to recognize the vast majority of target S-RNases. We found evidence of gene conversion events, which we suggest are essential to the constitution of a non-self recognition system and also contribute to self-compatible mutations.

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Supplementary Information for the manuscript

Gene duplication and genetic exchange drive the evolution of S-RNase based self-incompatibility

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This supplementary pdf file contains the following information:

Supplementary Methods

Supplementary Material

Supplementary Figures 1–13

Supplementary Tables 1–14

Supplementary Reference

Supplementary Methods

Plant materials. Lines of S_5 -, S_7 -, S_9 - and S_{11} -haplotypes from *Petunia hybrida*^{9,51} and the lines of S_{17} -, S_{19} - and S_{C2} -haplotypes from *P. axillaris*^{21,52} were described previously. In this research, we used additional lines of S_{10} -, S_{22} -, S_{0m} -, S_m -, and S_{22m} -haplotypes from *P. hybrida*. Homozygous self-compatible lines of S_m - and S_{0m} -haplotypes were the doubled-haploid line “Mitchell” and the inbred line W138, respectively⁵³. *S-RNase* sequences were identified by degenerate PCR using primer sequences listed in Supplementary Table 3. *S-RNase* sequence from Mitchell (S_m -*RNase*) was novel, but *S-RNase* from W138 was identical with previously identified S_0 -*RNase* from commercial SC *P. hybrida* cv. Strawberry Daddy⁵⁴. Because we cannot judge the identity of *S*-haplotype only from *S-RNase* sequence, we named *S*-haplotype from W138 as S_{0m} in this work. Homozygous lines of S_{10} -, S_{22} - and S_{22m} -haplotypes were established in this work from three commercial lines of *P. hybrida*. We determined S_{22m} - and S_{C2} -haplotypes as pollen-side SC haplotypes by reciprocal pollination with S_{22} - and S_{17} -homozygous plants, respectively (Supplementary Fig. 10a,b; see main text). We confirmed that SC phenotypes were genetically linked to the *S*-locus in all SC lines.

The interspecific crosses between the lines from *P. hybrida* and *P. axillaris* were compatible, and all the resulting progenies were fertile. Therefore, we do not distinguish them, and simply refer to them as ‘*Petunia*’ in this article.

Expression profiling. Template preparations and reaction conditions were described previously⁹. We conducted RT-PCR using *SLF* genes-specific primers (listed in Supplementary table 2) with 32 cycles for all of *SLF*s except for S_5 -*SLF18*, for which reaction was conducted with 28 cycles.

Vector construction. Fragments containing open reading frames (ORFs) of S_5 -*SLF3*, S_7 -*SLF3*, S_{11} -*SLF3B*, S_7 -*SLF9A* and S_{11} -*SLF9* were amplified with forward primers containing *Bam*HI sites and reverse primers containing *Sac*I sites. *pRI909-LAT52-pro:TAP:S₁₁-SLF3-35Spro:AtFT-LAT52-pro:Venus* plasmid vector⁹ was digested with *Bam*HI and *Sac*I, and the S_{11} -*SLF3* coding region was replaced with these amplified fragments to yield vectors listed in Supplementary Fig. 3.

Plant transformation and *in vivo* function assay. *S₇-SLF1*-expressing plants with *S₂₂*-haplotype were obtained by crossing with *S₇-SLF1*-transgenic plants previously described⁹. Other transgenic plants were newly obtained by *Agrobacterium*-mediated plant transformations described previously⁹. Breeding histories, genotypes and SI phenotypes of all transgenic lines are listed in Supplementary Table 4.

Genotyping and linkage analysis by genomic PCR. Genotyping of transgenic plants and linkage analysis were performed as described⁹. Progenies segregating for *S₅*-, *S₇*-, *S₉*-, *S₁₁*-haplotypes were obtained from *S₅S₁₁* × *S₇S₉* and *S₅S₉* × *S₇S₁₁* crosses, and progenies segregating for *S₁₇*- and *S₁₉*-haplotypes were obtained from *S₁₇S₁₉* × *S₅S₅* and *S₁₁S₁₁* × *S₁₇S₁₉* crosses. Gene-specific primers not described⁹ are listed in Supplementary Table 2.

Preparation of cDNA libraries and next generation sequencing. Total RNA was extracted from mature pollen of an *S₇*-homozygous plant and unopened mature anthers of *S₅*-, *S₉*-, *S₁₁*-, *S₁₇*-, *S₁₉*- and *S_{0m}*-homozygous plants, as described previously⁹. For sequencing of anther transcripts from *S_{0m}*-homozygous plant, poly (A)⁺ RNA were enriched using Oligotex-dT30 super (Takara), and the preparation of cDNA library and the sequencing using GS Junior system were performed at Roche Diagnostics Japan (<http://www.roche-diagnostics.jp>). For sequencing of transcripts from other plants, cDNA libraries were prepared from total RNA using the SMARTer PCR cDNA synthesis kit (Clontech). Sequencings were performed using Roche 454 Genome Sequencer FLX System at Hokkaido System Science (for *S₇*-homozygous plant), and at the Functional Genomics Center Zurich (for all other plants). The sequenced reads information is summarized in Supplementary Table 14.

Monte-Carlo simulation of S-RNase proportion recognized by SLFs. Monte-Carlo simulation was conducted in order to consider the difference of recognition rates among SLF types. The proportion of S-RNases recognized by *n* SLF types, $P_S(n)$, was simulated by the following equation (4):

$$P_S(n) = \frac{\sum_{i=1}^m P_{Mi}(n)}{m} \quad (4)$$

$$P_{Mi}(n) = 1 - \prod_{j=1}^n (1 - P_{Tj})$$

where m is the number of bootstrapping sampling times and the n elements of $\{P_{Tj}\}$ are selected in each bootstrapping sampling time i at random with replacement from the recognition rates of each SLF type,

$$P_T := \left\{ \frac{6}{12}, \frac{3}{6}, \frac{1}{6}, \frac{1}{8}, \frac{1}{8}, \frac{1}{8}, \frac{1}{8}, \frac{1}{6}, \frac{0}{6} \right\} \quad (5)$$

P_{Ti} corresponds to the recognition rate of SLF type i (Supplementary Table 13b). To calculate P_T , the recognition rates of SLF type, the interaction between SLF type and a particular S-RNase allelic variant is considered positive, when one or more allelic variants of the SLF type showed positive interaction experimentally. In this simulation, we used $m = 10,000$ times as bootstrapping sampling iteration. The Monte-Carlo simulation is implemented and performed in Ruby (ver. 1.9.3) (Supplementary Material). Mersenne Twister⁵⁵ was used for the random number generator in the Ruby script.

Supplementary Material Ruby script code that realizes the Monte-Carlo simulation for the estimation of the proportion of S-RNases recognized by n SLF types described in the Online Method.

```
#!/usr/bin/env ruby
# encoding: utf-8
# Date: 20140501
# Author: Masaomi Hatakeyama

TRIAL = 10000
N_MAX = 30
PROBS = [0.5, 0.5, 0.166666667, 0.125, 0.125, 0.125, 0.125, 0.166666667, 0.0]
SEED = 1234

srand(SEED)
dat_file = File.basename(__FILE__).gsub(/.rb/, '.dat')
plt_file = File.basename(__FILE__).gsub(/.rb/, '.plt')
png_file = File.basename(__FILE__).gsub(/.rb/, '.png')

class Array
  def sum
    inject(0.0) { |sum, i| sum += i }
  end
  def ave
    inject(0.0) { |sum, i| sum += i } / size
  end
  def var
    average = ave
    inject(0.0) { |sum, i| sum += (i - average)**2 } / (size-1)
  end
  def sd
    Math::sqrt(var)
  end
end
```

```

def se
  sd/Math.sqrt(size)
end
def ci
  1.96*se
end
end

def f(n,probs)
  fail_prob = 1.0
  n.times do
    px = probs[rand(probs.length)]
    fail_prob *= (1.0-px)
  end
  1.0-fail_prob
end

open(dat_file, "w") do |out|
  (1..N_MAX).each do |n|
    trials = []
    trials2 = []
    TRIAL.times do
      trials << f(n, PROBS)
    end
    #out.puts [n, trials.ave, trials.sd].join("¥t")
    out.puts [n, trials.ave, trials.ave-trials.sd, trials.ave+trials.sd].join("¥t")
  end
end

open(plt_file, "w") do |out|
  out.print <<-EOF
  set xlabel "Number of SLF types"
  set ylabel "SRNase recognized probability by SLF types"

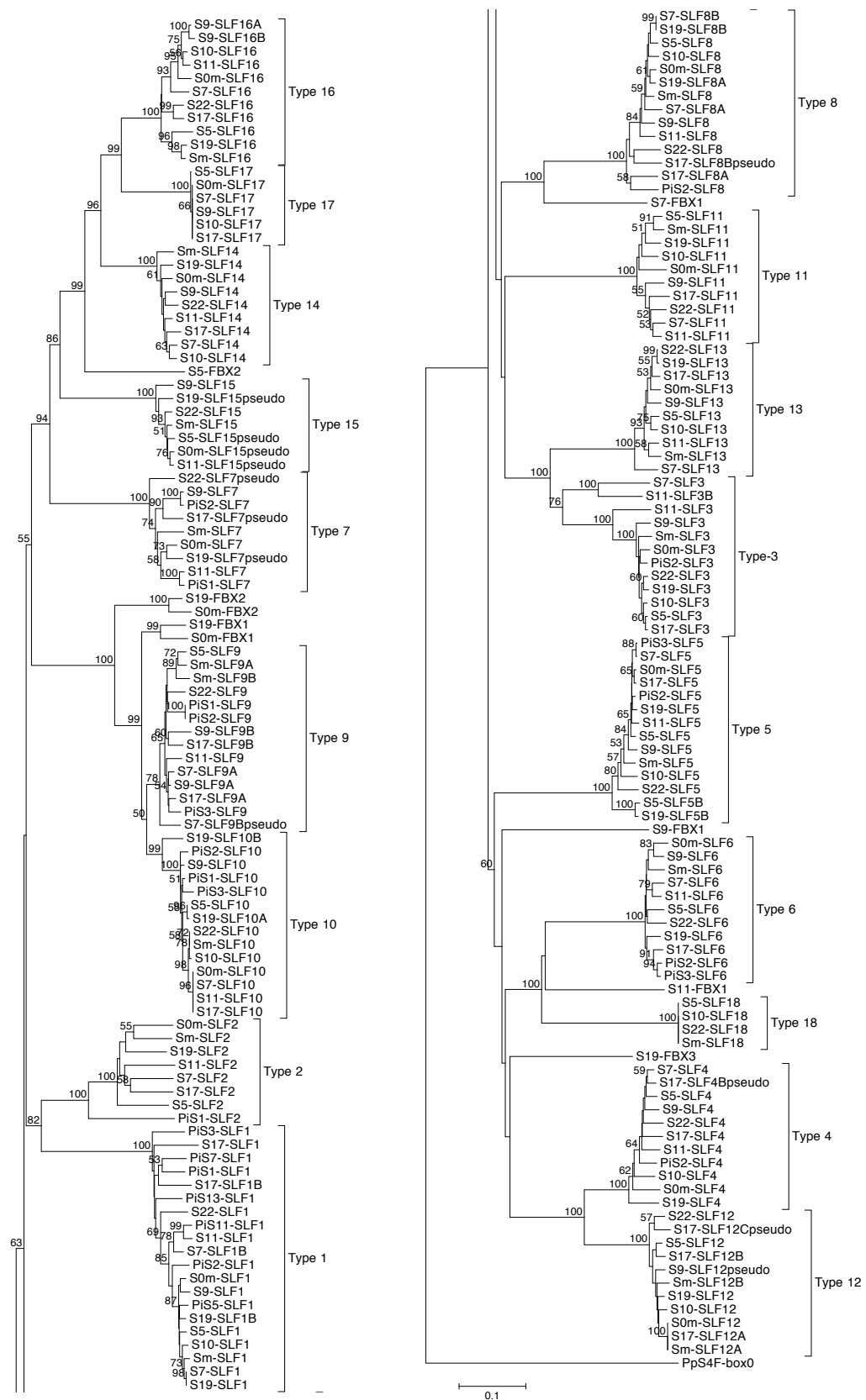
```

```

set yrange[0.1:1.0]
set key below
set term png
set out "#{png_file}"
set arrow 1 from 20,0.1 to 20,1 nohead linewidth 0.1
set arrow 2 from 16,0.1 to 16,1 nohead linewidth 0.1
set arrow 3 from 18,0.1 to 18,1 nohead linewidth 0.1
set xtics('0' 0, '5' 5, '10' 10, '15' 15, '16' 16, '18' 18, '20' 20, '25' 25, '30' 30)
p "#{dat_file}" not w l, "#{dat_file}" u 1:2:3:4 not with yerrorbar, 0.95 not, 0.99 not
    EOF
end

command = "gnuplot -persist #{plt_file}"
puts command
system command

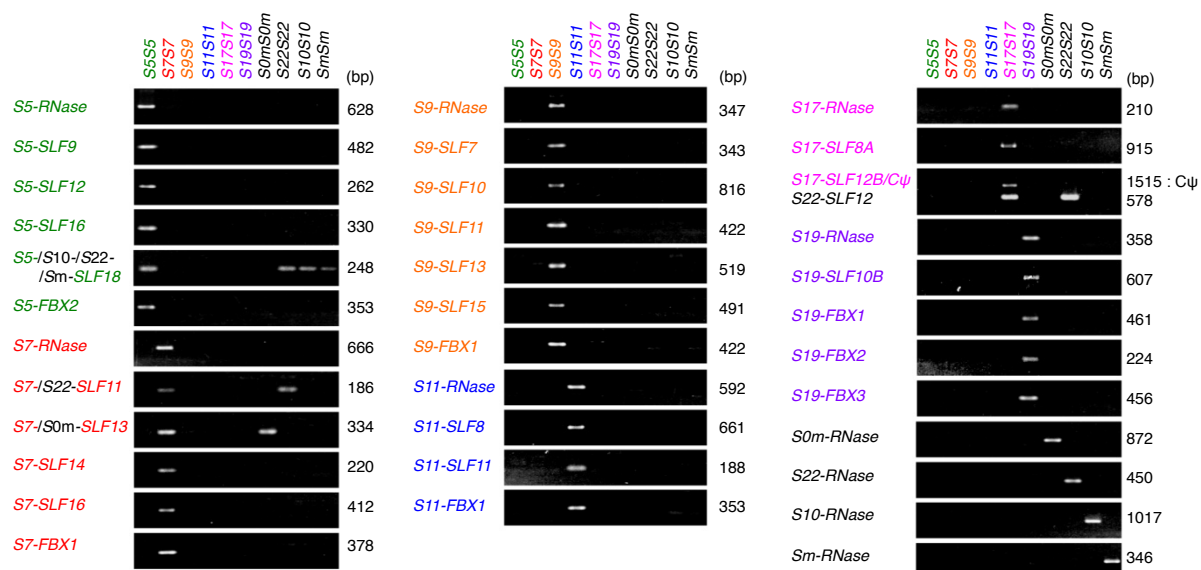
```



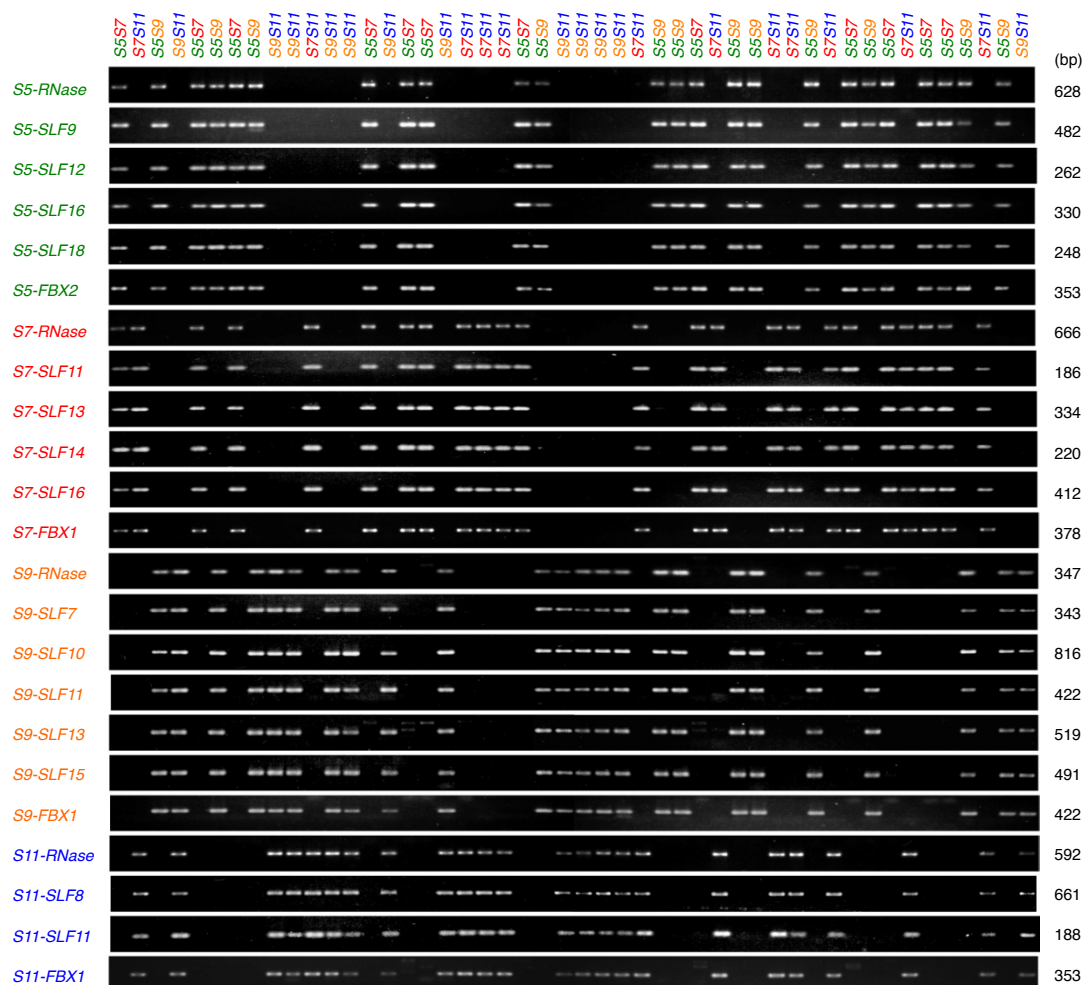
sup. Fig. 1

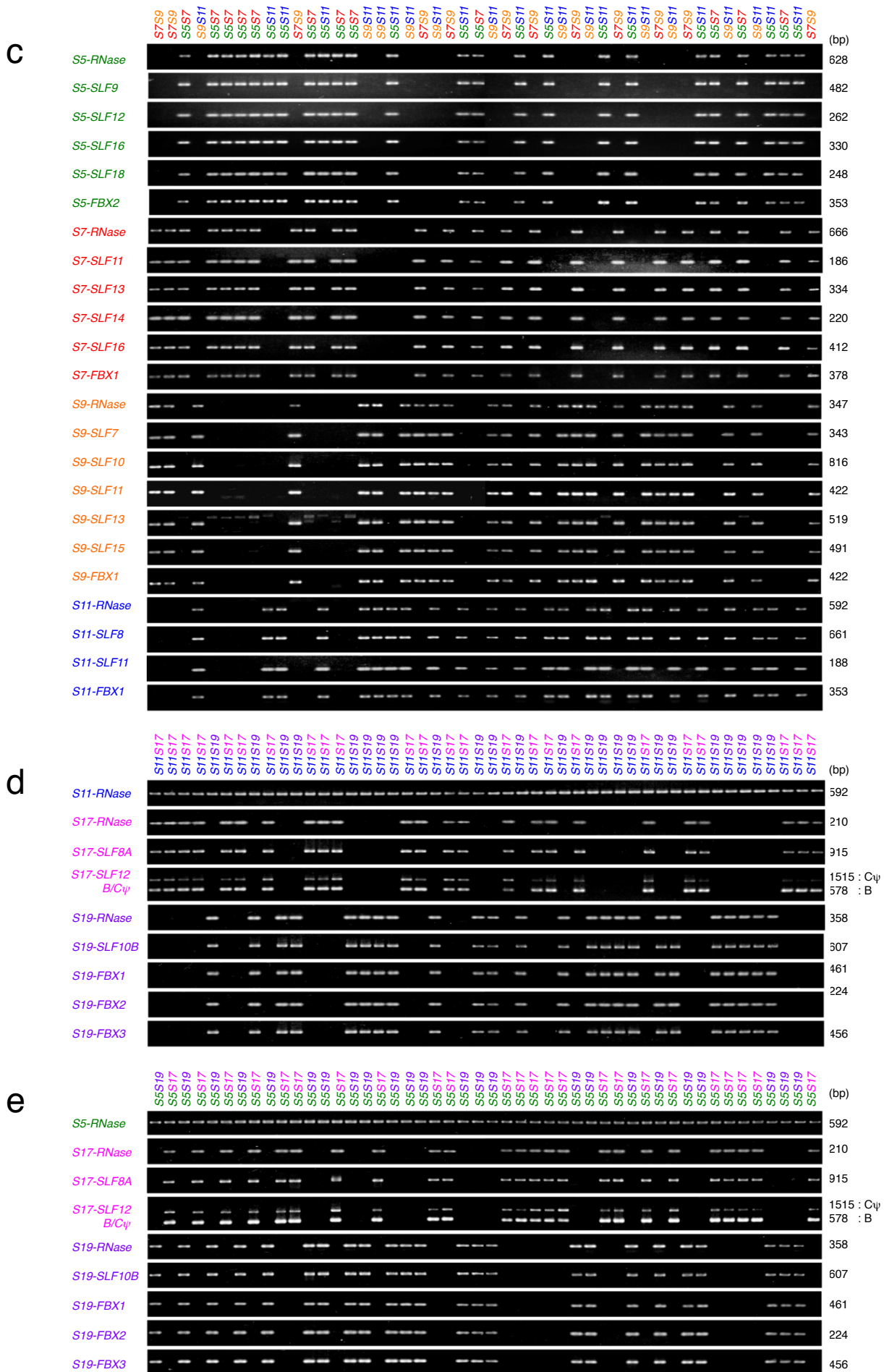
Supplementary Figure 1 Phylogeny of SLFs from *Petunia*. Phylogenetic tree of deduced amino-acid sequences of SLFs shown in Fig.1 is described without the compressed representation. For details, see METHODS and caption of Fig. 1. Based on this phylogeny, previously identified *S₇-SLF3* (Ref. 9) is now assigned into a clade type-13, and renamed as *S₇-SLF13*. Instead, we newly identified *SLF* that belongs to the type-3 clade from *S₇*-haplotype, and named it as *S₇-SLF3*. The phylogeny also suggests that the previously identified *SLF*-like genes, *SLFLa*, *SLFLb*, *A113* and *A134*, are alleles of type 7–10 *SLFs*, and renamed as *SLF7*, *SLF8*, *SLF9*, and *SLF10*, respectively^{37,56}. Information of accession codes and synonyms of *SLFs* used in this phylogenetic analysis was summarized in Supplementary Tables 1 and 9.

a



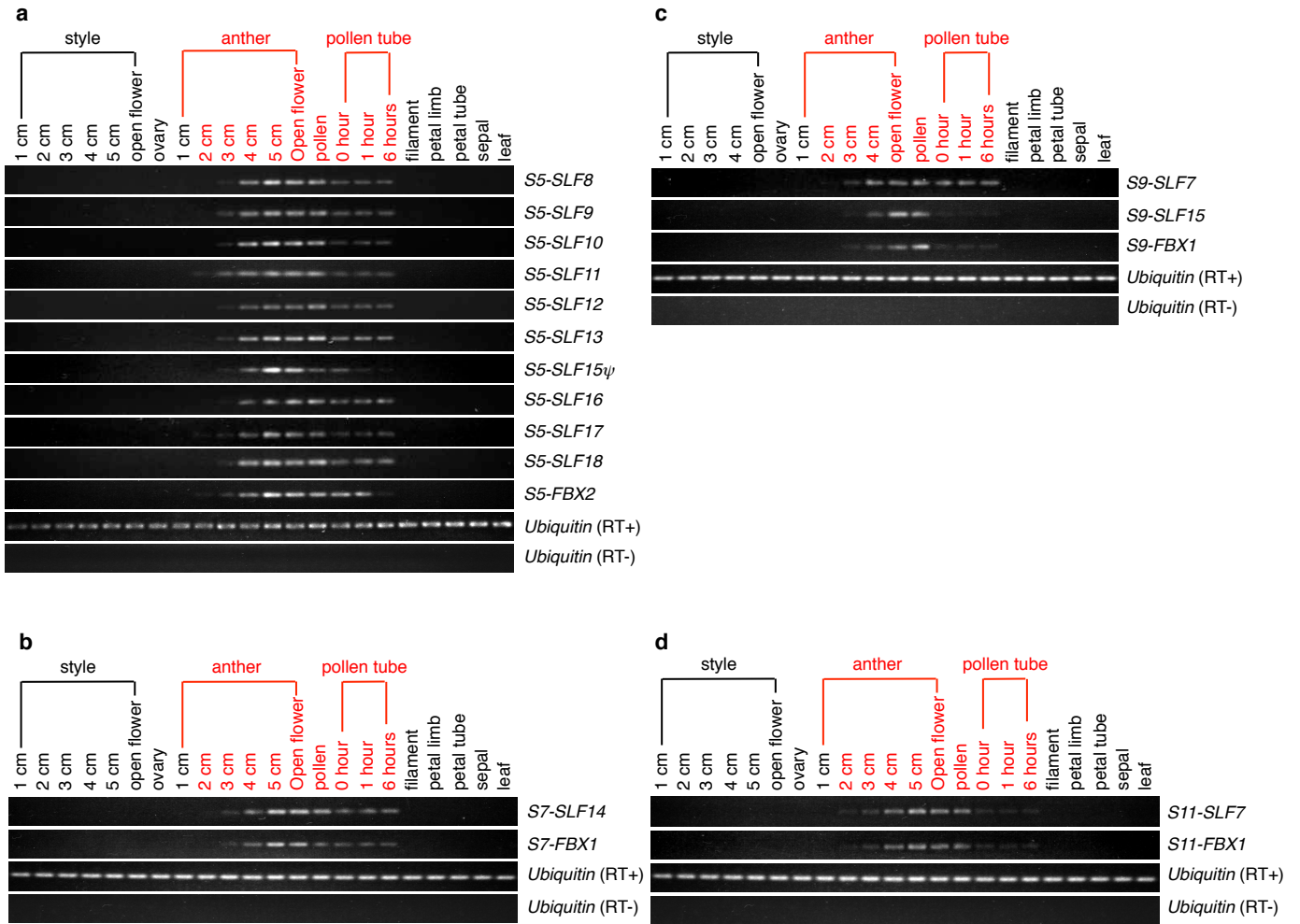
b





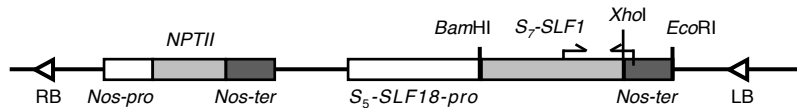
sup. Fig. 2

Supplementary Figure 2 Genetic linkage between *SLF*s and *S-RNase*. **(a)** PCR was performed using genomic DNA prepared from *S*-homozygous plants to evaluate the specificity of primer pairs. Primers are listed in Supplementary Table 2. **(b, c)** Genetic linkage analyses between *S-RNase* and *SLF* genes of *S*₅-, *S*₇-, *S*₉- and *S*₁₁-haplotypes. PCR was performed using genomic DNA prepared from 48 progeny plants of an *S*₅*S*₁₁ × *S*₇*S*₉ cross (b), and from 48 progeny plants of an *S*₅*S*₉ × *S*₇*S*₁₁ cross (c). **(d, e)** Genetic linkage analyses between *S-RNase* and *SLF* genes of *S*₁₇- and *S*₁₉-haplotypes. PCR was performed on genomic DNA prepared from 48 progeny plants of an *S*₁₁*S*₁₁ × *S*₁₇*S*₁₉ cross (d) and from 48 progeny plants of an *S*₁₇*S*₁₉ × *S*₅*S*₅ cross (e).

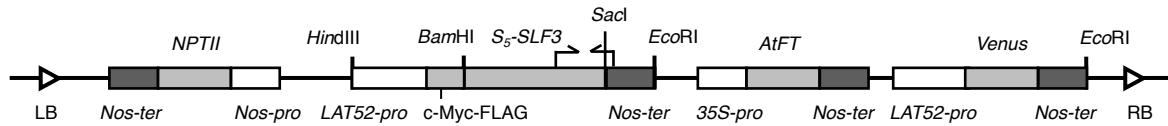


Supplementary Figure 3 Newly isolated *SLF* genes show a male reproductive organ-specific expression profile. RT-PCR analyses of the expression profiles of six types of *SLF* genes in (A) *S*₅-, (B) *S*₇-, (C) *S*₉- and (D) *S*₁₁-homozygotes. cDNA was prepared from each *S*-homozygote, and Type-7 to Type-18 *SLFs* and ungrouped *SLF*-like F-box (*FBX*) genes were amplified by specific primer pairs. The *ubiquitin* gene was also amplified and electrophoresed as a control. All primer sequences are shown in Supplementary Table 2.

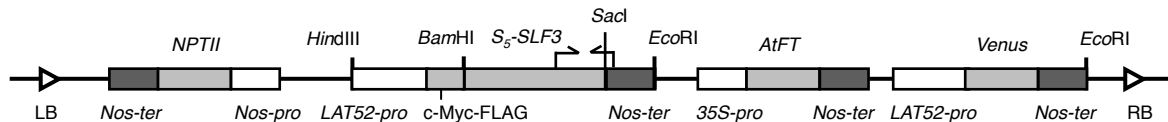
pBI121-S₅-FBX-pro:S₇-SLF1



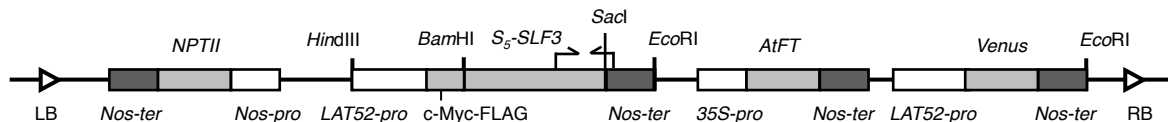
pRI909-LAT52-pro:TAP:S₅-SLF3-35S-pro:AtFT-LAT52-pro:Venus



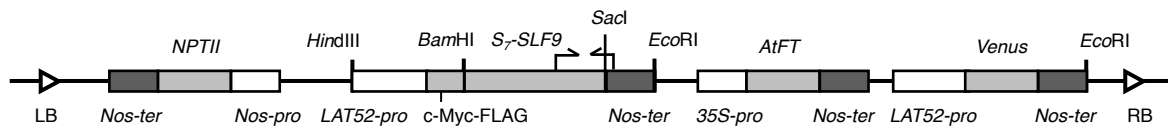
pRI909-LAT52-pro:TAP:S₇-SLF3-35S-pro:AtFT-LAT52-pro:Venus



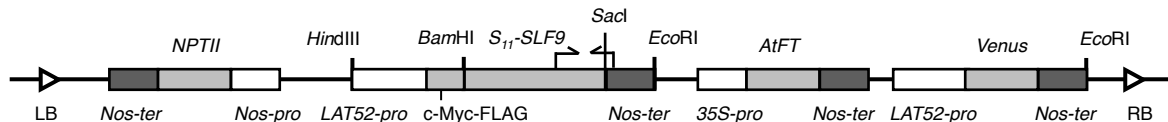
pRI909-LAT52-pro:TAP:S₁₁-SLF3B-35S-pro:AtFT-LAT52-pro:Venus



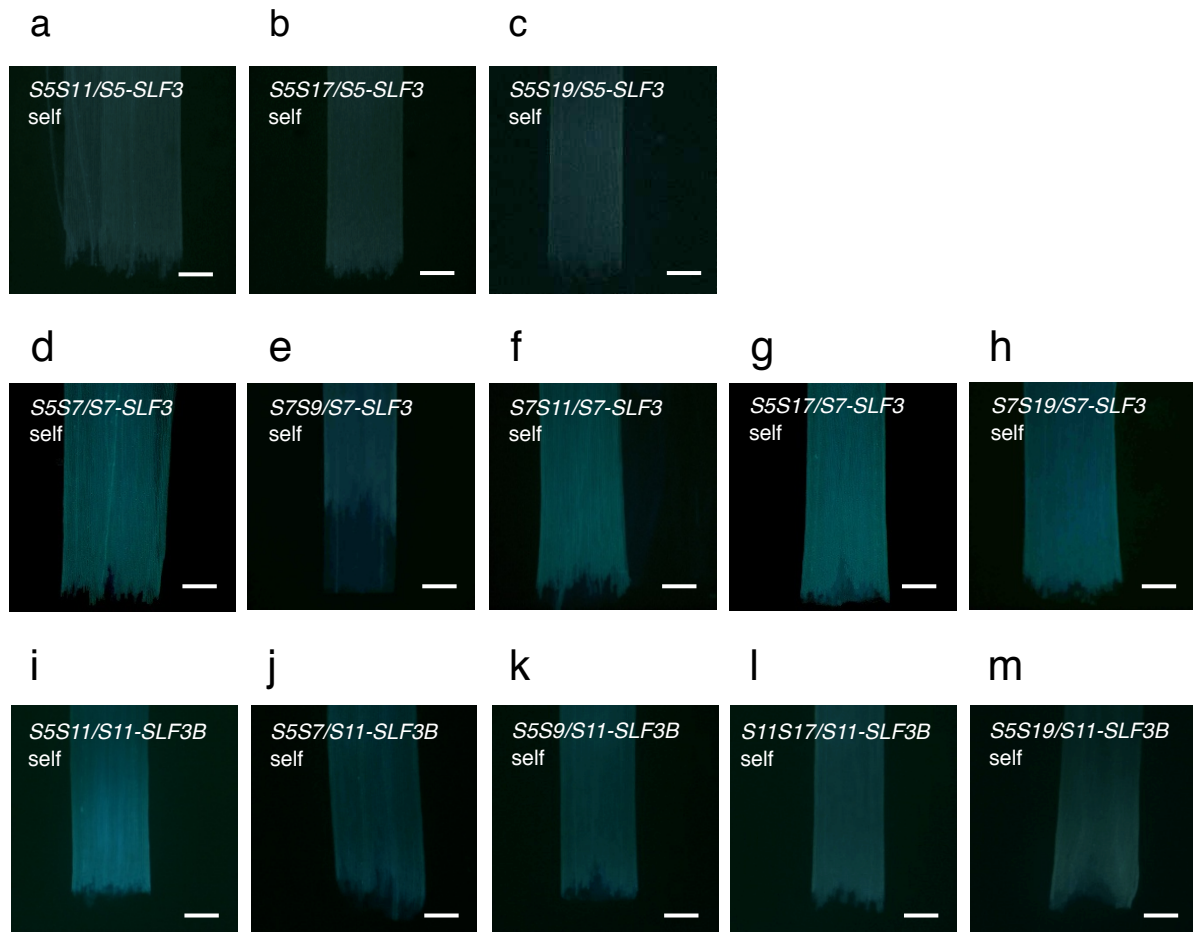
pRI909-LAT52-pro:TAP:S₇-SLF9A-35S-pro:AtFT-LAT52-pro:Venus



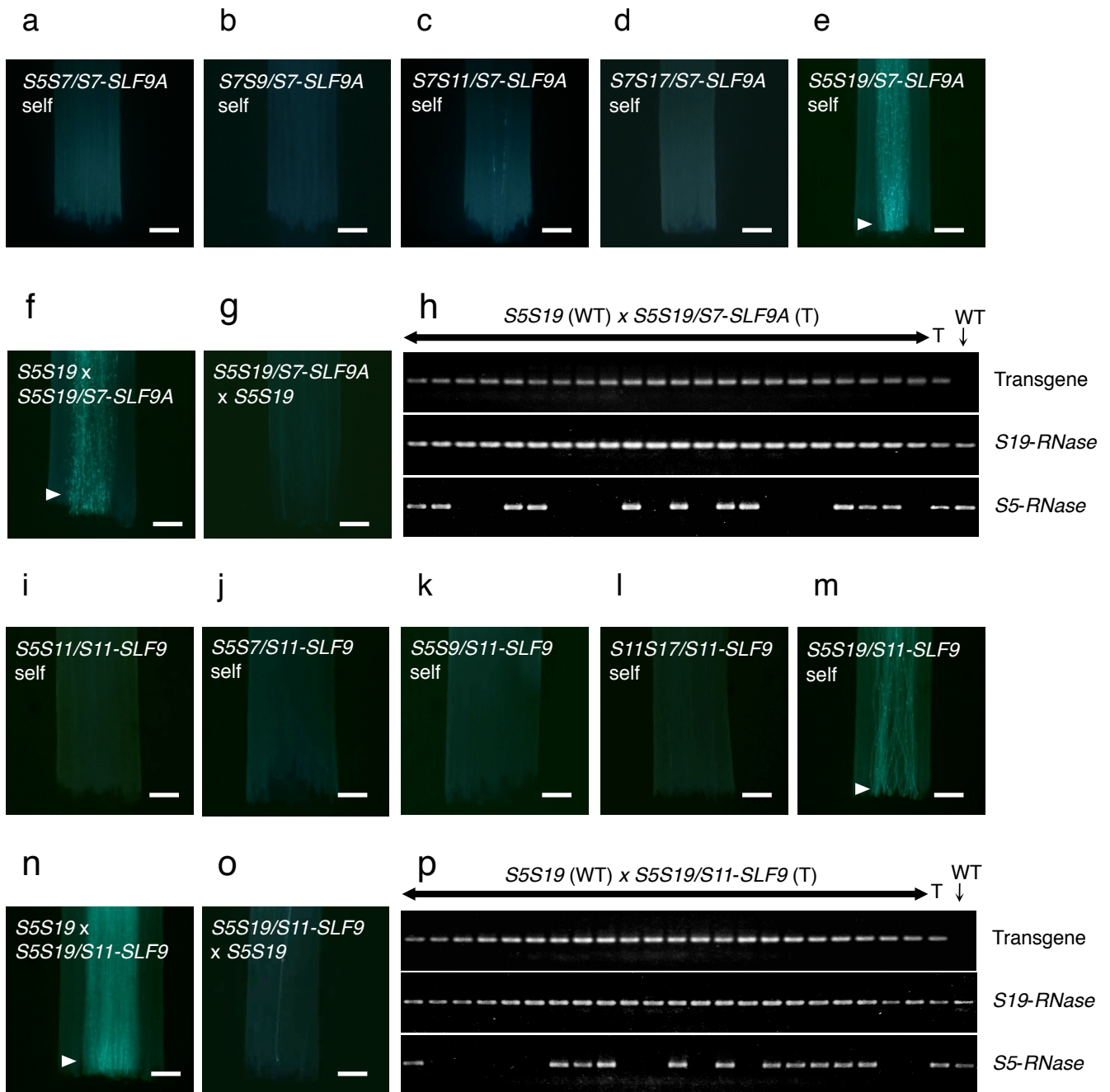
pRI909-LAT52-pro:TAP:S₁₁-SLF9-35S-pro:AtFT-LAT52-pro:Venus



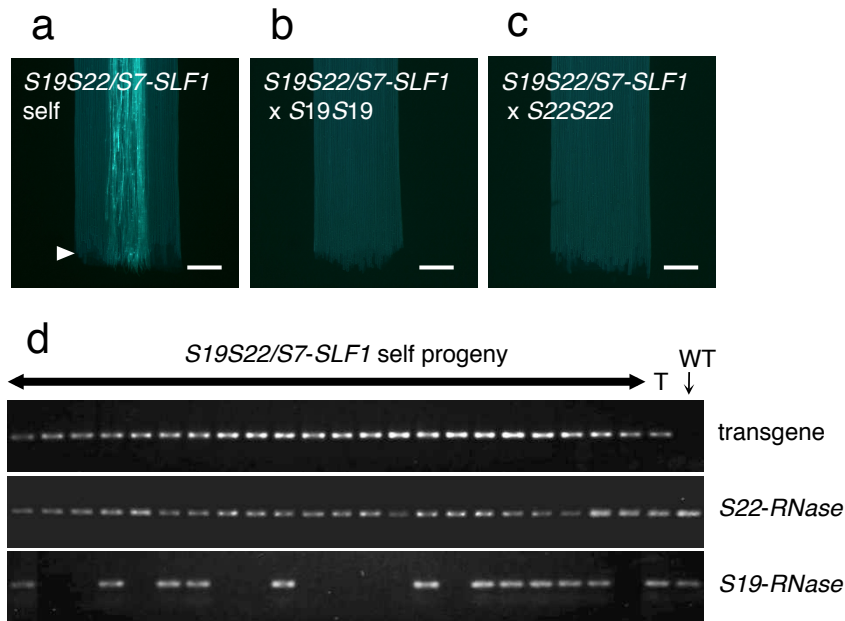
Supplementary Figure 4 Schematic representation of transgene constructs used for transformation experiments. The region between the right border (RB) and left border (LB) is integrated into transgenic plants. *Nos*, the gene encoding nopaline synthase; *pro*, promoter; *ter*, transcription terminator; *LAT52-pro*, pollen/pollen tube-specific *LAT52* promoter of tomato⁵⁷; *35S-pro*, Cauliflower mosaic virus 35S promoter; *S₅-SLF18-pro*, 2028 bp of the upstream region of the *S₅-SLF18* gene; *NPTII*, the gene encoding neomycin phosphotransferase II (conferring kanamycin resistance); *AtFT*, *Arabidopsis thaliana* *FLOWERING LOCUS T* gene (accelerating flowering and to shortening the experimental time period)⁵⁸; *Venus*, the gene encoding enhanced yellow fluorescent protein⁵⁹. Small arrows denote the locations of the forward and reverse primers used in PCR genotyping and RT-PCR analysis of each transgene (for primer sequences, see Supplementary Table 2).



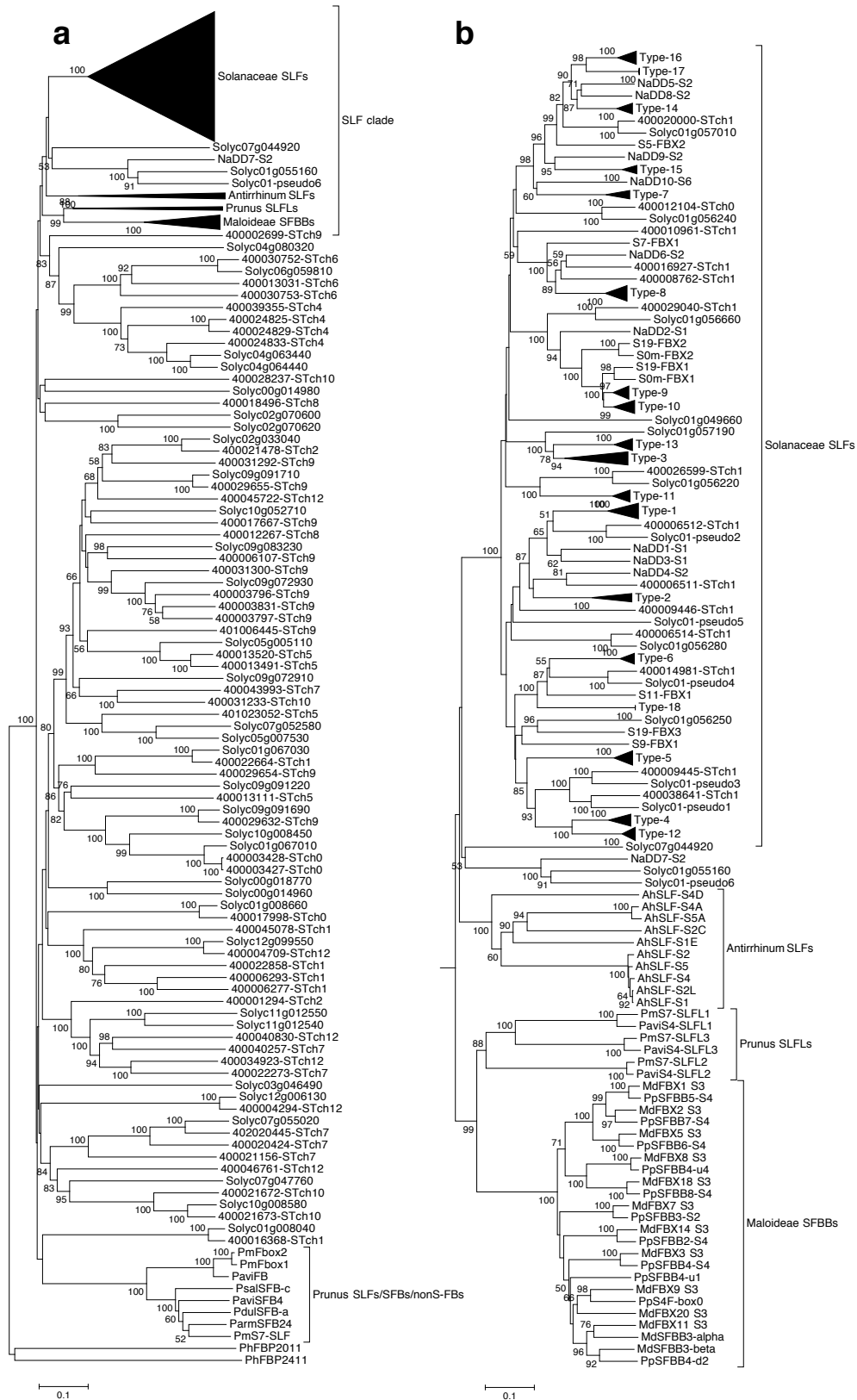
Supplementary Figure 5 Representative results of transgenic experiments to detect the *in vivo* interaction between type-3 SLFs and S₇-RNase. **(a–c)** Pollen tubes derived from S₅-SLF3 transgenic plants were stained with aniline blue and monitored by fluorescence microscopy. The S₅S₁₁/S₅-SLF3 **(a)**, S₅S₁₇/S₅-SLF3 **(b)** and S₅S₁₉/S₅-SLF3 **(c)** transgenic plants retained SI, indicating that S₅-SLF3 did not recognize and detoxify S₅-, S₁₁-, S₁₇-, and S₁₉-RNases. **(d–h)** Pollen tubes derived from S₇-SLF3-transgenic plants were stained with aniline blue and monitored by fluorescence microscopy. All transgenic plants exhibited SI, indicating that S₇-SLF3 did not recognize and detoxify S₅-, S₇-, S₉-, S₁₁-, S₁₇-, and S₁₉-RNases. **(i–m)** Pollen tubes derived from S₁₁-SLF3B-transgenic plants were stained with aniline blue and monitored by fluorescence microscopy. All transgenic plants exhibited SI, indicating that S₁₁-SLF3B did not recognize and detoxify S₅-, S₇-, S₉-, S₁₁-, S₁₇-, and S₁₉-RNases. Bars = 200 μm.



Supplementary Figure 6 Representative results of transgenic experiments to demonstrate *in vivo* interaction between type-9 SLFs (S_7 -SLF9A, and S_{11} -SLF9) and S_{19} -RNase. **(a–g)** Pollen tube growth derived from S_7 -SLF9A-transgenic plants. S_5S_7/S_7 -SLF9A **(a)**, S_7S_9/S_7 -SLF9A **(b)**, S_7S_{11}/S_7 -SLF9A **(c)** and S_7S_{17}/S_7 -SLF9A **(d)** retained SI, suggesting that S_7 -SLF9A does not recognize S_5 -, S_7 -, S_9 -, S_{11} - and S_{17} -RNases. S_5S_{19}/S_7 -SLF9A **(e)** exhibited SC. The reciprocal cross **(f, g)** indicated the SI breakdown occurred in pollen. **(h)** PCR genotyping of 22 progeny plants obtained by crossing S_5S_{19} (WT) with pollen from S_5S_{19}/S_7 -SLF9A (T). These results suggest that S_7 -SLF9A recognizes S_{19} -RNase as a target. **(i–p)** Pollen tube growth derived from S_{11} -SLF9-transgenic plants. S_5S_{11}/S_{11} -SLF9 **(i)**, S_5S_7/S_{11} -SLF9 **(j)**, S_5S_9/S_{11} -SLF9 **(k)** and $S_{11}S_{17}/S_{11}$ -SLF9 **(l)** retained SI, suggesting that S_{11} -SLF9 does not recognize S_5 -, S_7 -, S_9 -, S_{11} - and S_{17} -RNases. S_5S_{19}/S_{11} -SLF9 **(m)** exhibited SC. The reciprocal cross **(n, o)** indicated the SI breakdown occurred in pollen. **(p)** PCR genotyping of 22 progeny plants obtained by crossing S_5S_{19} (WT) with pollen from S_5S_{19}/S_{11} -SLF9 (T). These results suggest that S_{11} -SLF9 recognizes S_{19} -RNase as a target. Bars = 200 μ m.



Supplementary Figure 7 Representative results of transgenic experiments to demonstrate the *in vivo* interaction between *S7-SLF1* and *S22-RNase*. **(a-c)** Pollination phenotype of *S19S22/S7-SLF1* transgenic plants. *S19S22/S7-SLF1* exhibited breakdown of SI (a), whereas pistil of *S19S22/S7-SLF1* retained incompatibility against pollen from *S19*-homozygote (b) and *S22*-homozygote (c). **(d)** PCR genotyping of 22 progeny plants obtained by selfing of *S19S22/S7-SLF1* (T).

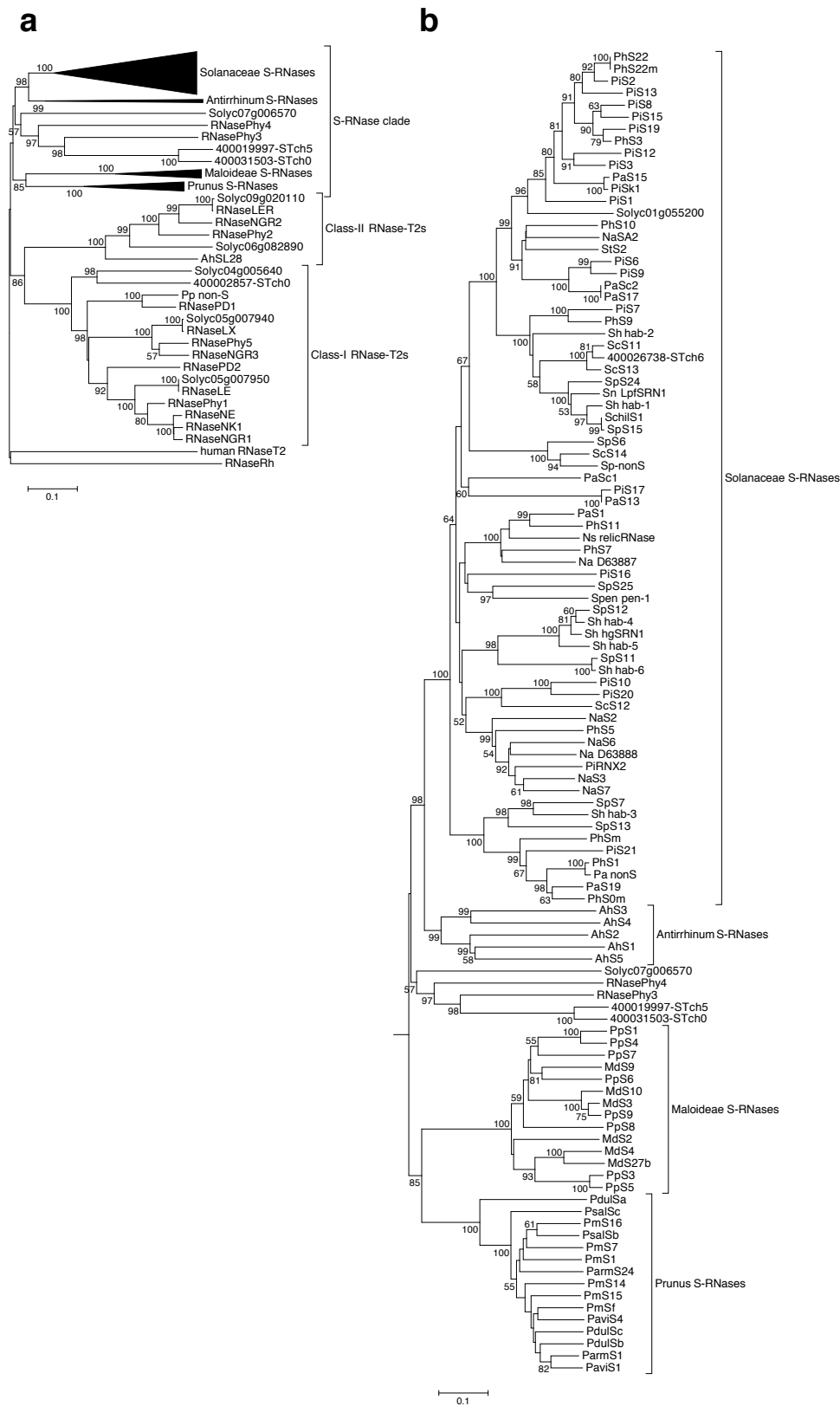


sup. Fig. 8

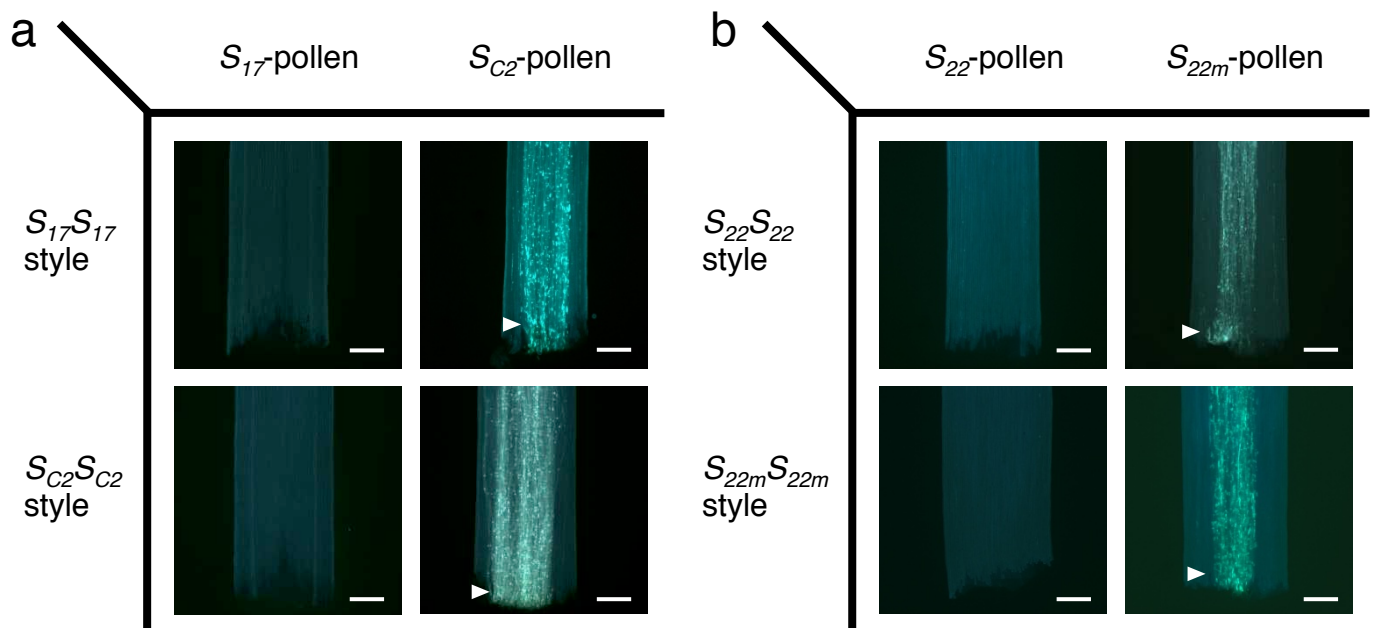


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Supplementary Figure 8 Phylogenies of SLF-related F-boxes from species possessing S-RNase-based SI. **(a)** Whole tree constructed by using *Petunia* SLFs and all BLAST hits from the tomato and potato databases, as well as other SLF-related F-box proteins from Solanaceae, Plantaginaceae, and Rosaceae. PiFBP2011 and PiFBP2411 from *Petunia inflata* were used as the outgroup⁶⁰. For other details, see Online Methods and the caption of Fig. 3. In this tree, Solanaceae SLF, Antirrhinum SLF, Prunus SLFL, and Maloideae SFBB clades are shown in compressed representation (black triangles). A subtree of Solanaceae SLFs is shown in Fig. 3a. **(b)** Subtree of SLF clade in (a) is shown. *Petunia* SLFs types are shown in compressed representation (black triangles). **(c)** Subtree of Solanaceae SLF clade (see Fig. 3a) is shown without compression. Round red markers indicate *SLFs* from genus *Petunia* (*P. hybrida*, *P. axillaris* and *P. inflata*); blue, genus *Solanum* (*Solanum lycopersicum*, *S. tuberosum* and their relatives); green, genus *Nicotiana*. 'PGSC0003DMG' is omitted from each potato gene ID to simplify, and '-STchx' is attached to indicate genes located on *S. tuberosum* chromosome x.



Supplementary Figure 9 Phylogenies of S-RNase-related RNase-T2s from species possessing S-RNase-based SI. **(a)** Whole tree constructed using *Petunia* S-RNases and all hits from BLAST searches of the tomato and potato databases, as well as other RNase-T2s from Solanaceae, Plantaginaceae and Rosaceae. Classification of RNase-T2 is made according to refs. 61 and 62. RNase-T2s from human and the filamentous fungus *Rhizopus niveus* were used as outgroup^{63,64}. For other details, see METHODS and the caption of Fig. 3. In this tree, S-RNase clades from Solanaceae, Antirrhinum, Prunus, and Maloideae are shown in compressed representation (black triangles). A subtree of Solanaceae S-RNases clade is shown in Fig. 3b. **(b)** Subtree of S-RNase clade in (a) is shown without compression.



Supplementary Figure 10 S_{C2-} and S_{22m} -haplotypes are the pollen-side SC haplotypes. Reciprocal pollination tests between SC S_{C2-} and SI S_{17} -homozygous plants (a) and between SC S_{22m-} and SI S_{22} -homozygous plants (b). Pollen tubes were stained with aniline blue, and the compatibility was judged by the presence of a large number of pollen tubes at the basal end of the style (arrowhead). Bars = 200 μ m.

a. S-RNases

S7-RNase	1	MFRSGLNSASFLFLAQNPGYGVFDQTLVLTWPPSPFCHTRP-CRR--TPRNFTHGLWPDQHVLIINDCKDT--YTTISDAREKEDARNDPLRYTER	95
S19-RNase	1	KL-SLVY-SLF-PS-E-S-HW...AGY-KL-G-P-FVIA-N...NVKTM-N...PSTRFNK-AV-EQ-N...EK...E-PSLAH	99
SC2-RNase	1	K-L-H-H-LF-DS-E-D-HM...AA-YF-GF-O-IF-R...KERQR-QF-A-DYK-VNF-EGDI-SS-HH-IQ-LFNKE	98
S22m-RNase	1	L-L-H-H-LF-DS-SAN-DF...A-YF-NK-O--RSN...EKK-FR-EP-PGD-KFSRFKEDNIINV-ERH-IQM-PD-D	97
S7-RNase	96	DAHQLOSTWRYEYAKHGGCCSERFDQSLVFNLANAKDKSEHLLQMLIOGHPGKTPVDKSEFANVTH-EYDNECQGGPYKMLLEIGICNDPFA	194
S19-RNase	100	IGTST-L-K-L-E-FPA-S-S-DF-IR...FD-T-L-S-V-F-S-IGER-NSS-AS-R-VK...E-L-YYOC...E-F-RTT	197
SC2-RNase	99	VGLKY-PL-HDG-EL...-NL-E-F-MR-S-D-SL-HM...T-RR-LFOR-EGC-L-Q-MD-S-K-BHIGV-L...F-L-N	197
S22m-RNase	98	YNAK-PL-QH-L-R-E-KNL...-N-L-IR...FD-T-L-IF-T...FKHP-GE-OR-L-L-NNKD-S-K-A-ENIKGV-L...F-A	197
S7-RNase	195	TKVTFCHR-RKTKPLNKKETSPQ	218
S19-RNase	198	VAMMS-P-LSTG-FGTNAR-L-R	222
SC2-RNase	198	ESPYH-PQ-SHS-EKRGYTC-L-R	220
S22m-RNase	198	DSFHD-RR-SY-DETSTQTLLRR	221

b. SLF1s

S7-SLF1	1	MANGILKKLPEDLVFLILLTPFKSLLRFCISKANSILIQSTTFINRHINKNTNTKAEFIFLKRSIKDEEEFINILSFFSGDDVLNPLFPDIDVSYM	100
S19-SLF1	1		100
SC2-SLF1C	1		100
S22m-SLF1B	1		100
S7-SLF1B	1		100
S19-SLF1B	1		100
SC2-SLF1A	1		100
SC2-SLF1B	1		100
S22m-SLF1A	1		100
S7-SLF1	101	TSKCDCTPTPLIGPCDGLIALDITITVINPATRNFRVLPPSPFGCPKGYHRSVEGVGFGPDTISWYKVVRISEVYCEADGYPGKDKSIDVCDLST	200
S19-SLF1	101		200
SC2-SLF1C	101		200
S22m-SLF1B	101		200
S7-SLF1B	101		200
S19-SLF1B	101		200
SC2-SLF1A	101		200
SC2-SLF1B	101		200
S22m-SLF1A	101		200
S7-SLF1	201	DSWRELDHVQLPSIYVWPCAGMLYKEMVHWFATTTDSMVILCFDPMSTEMFHDMKMPDTCRSRITHELYYGLVILCESFTLIGYSNPISIDPVEDKMHIV	300
S19-SLF1	201		300
SC2-SLF1C	201		300
S22m-SLF1B	201		300
S7-SLF1B	201		299
S19-SLF1B	201		300
SC2-SLF1A	201		299
SC2-SLF1B	201		300
S22m-SLF1A	201		299
S7-SLF1	301	MMYGVSESWIMKYTIRPLSIESPLAVNKHILLQSRGILLISYDLNSGEARDLNLHGFPDLSVKVYKECLTSIPKGESEYTKVQRT	389
S19-SLF1	301		389
SC2-SLF1C	301		389
S22m-SLF1B	301		389
S7-SLF1B	301		388
S19-SLF1B	301		389
SC2-SLF1A	301		388
SC2-SLF1B	301		389
S22m-SLF1A	300		388

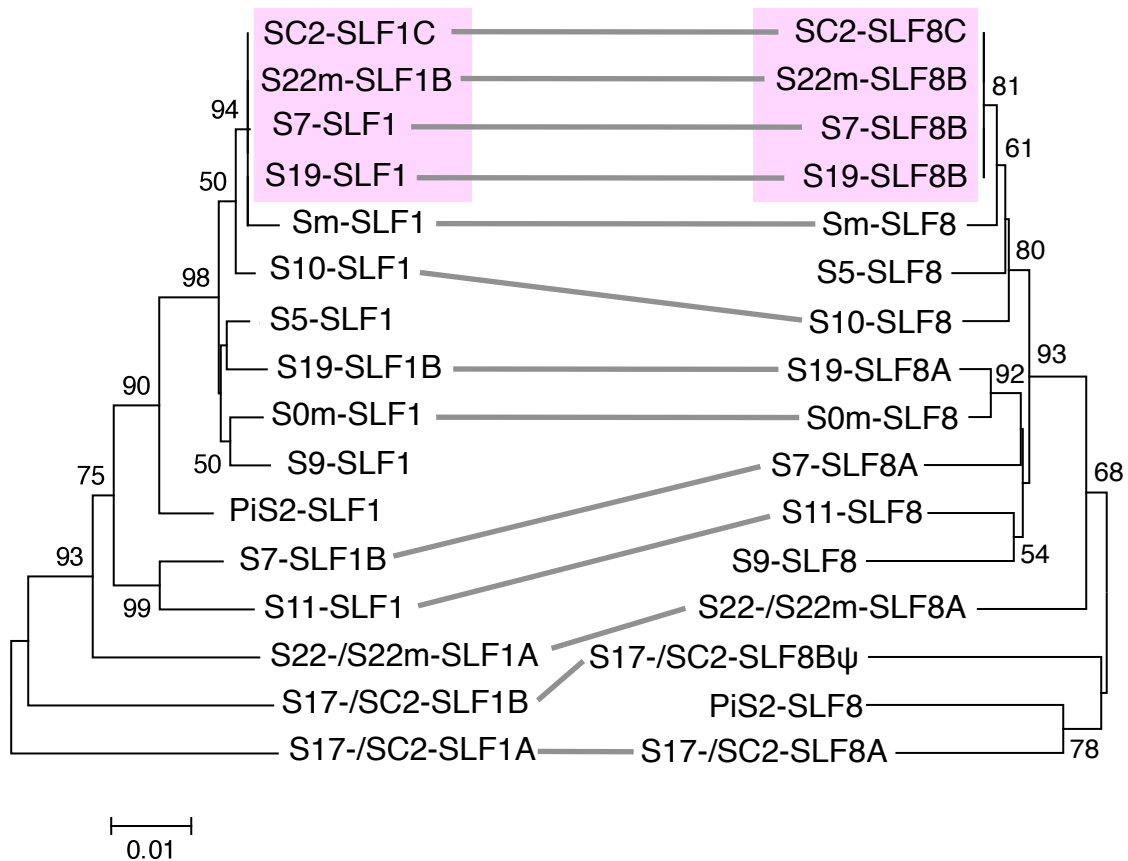
c. SLF8s

S7-SLF8B	1	MMLDGIMKHLPEDIAMYL-LRFPVKSLLRFKFKISKWSLTIESFTFINIHLNRATTTKNGFLFSRSYREETEGFKNVLSILSSGNNDDLIPVSDLSL	99
S19-SLF8B	1		99
SC2-SLF8C	1		99
S22m-SLF8B	1		99
S7-SLF8A	1		99
S19-SLF8A	1		99
SC2-SLF8A	1		99
SC2-SLF8B	1		99
S22m-SLF8A	1		98
S7-SLF8B	100	DYLTFTEYLLFNKLVGPCNGLVLTDFEIVFLFNPAKTNMILPPSPFVCPKGFHRSFRGGVGFGFDSIVKDYKFTTISEVFMDSWVPDEKQKVEVYD	199
S19-SLF8B	100		199
SC2-SLF8C	100		199
S22m-SLF8B	100		199
S7-SLF8A	100		199
S19-SLF8A	100		199
SC2-SLF8A	100		199
SC2-SLF8B	100		199
S22m-SLF8A	99		198
S7-SLF8B	200	LRFDSDWRDLNHVDQQLPTVYVYPCFEMLYNGAFHWYAINDRDLHVILSFDISTEIFSISKMPATGKSSGGKKYGLIVLNSLTLICYPNPDCEMDPSKDS	299
S19-SLF8B	200		299
SC2-SLF8C	200		299
S22m-SLF8B	200		299
S7-SLF8A	200		299
S19-SLF8A	200		299
SC2-SLF8A	200		295
SC2-SLF8B	200		299
S22m-SLF8A	199		298
S7-SLF8B	300	MDIWMMEYGHYESWTKKYIKPLPIESPLTIWRDHLILLQSKSGLLVSYDLSSNEVKEFDLHGYPKSLRVLVYKESLISIPKRGCKHGTFKNGKRGIT	399
S19-SLF8B	300		399
SC2-SLF8C	300		399
S22m-SLF8B	300		399
S7-SLF8A	300		391
S19-SLF8A	300		394
SC2-SLF8A	296		389
SC2-SLF8B	300		391
S22m-SLF8A	299		393
S7-SLF8B	400	ISY	402
S19-SLF8B	400	---	402
SC2-SLF8C	400	---	402
S22m-SLF8B	400	---	402
S7-SLF8A	391	---	391
S19-SLF8A	394	---	394
SC2-SLF8A	389	---	389
SC2-SLF8B	391	---	391
S22m-SLF8A	393	---	393

Supplementary Figure 11 S_7 -, S_{19} -, S_{22m} -, S_{C2} -haplotypes share identical *SLF1* and *SLF8*. Alignment of deduced amino-acid sequences of S-RNases (**a**), SLF1s (**b**) and SLF8s (**c**) from S_7 -, S_{19} -, S_{22m} -, and S_{C2} -haplotypes of *Petunia*. Deduced amino-acid sequences were aligned using MEGA5.2.2 and illustrated using the GENETYX-MAC (ver. 16.0.6). Amino-acid residues conserved in more than half of the aligned sequences are shown by darkened background. Amino-acid residues identical with the first line are indicated by dots. Gaps are indicated by hyphens.

Type-1 SLFs

Type-8 SLFs



Supplementary Figure 13 Coevolutionary relationships between type-1 and type-8 SLFs. Neighbor-joining phylogenetic trees of type-1 and -8 SLFs were created with MEGA 5.2.2. Both trees are shown in the same scale; the bar for each tree indicates the number of base substitutions per site. Numbers on the branches indicate bootstrap values >50% with 1,000 trials. Identical SLF1 and SLF8 shared among four different *S*-haplotypes are highlighted in pink. Horizontal gray lines indicated gene pairs on the same *S*-haplotypes holding similar branching patterns.

Supplementary Table 1 Repertoire of SLF genes identified in our research

S5-haplotype

Gene	RPKM	Accession ID	References	Note
<i>S5-SLF1</i>	23.72	AB568390	ref. 9	interact with S9- and S17-RNases
<i>S5-SLF2</i>	2.28	AB568394	ref. 9	interact with S9- and S11-RNases
<i>S5-SLF3</i>	13.50	AB568399	ref. 9	interact with S7-RNases
<i>S5-SLF4</i>	3.06	AB568405	ref. 9	
<i>S5-SLF5</i>	2.37	AB568411	ref. 9	
<i>S5-SLF5B</i>	10.19	AB932964	this work	
<i>S5-SLF6</i>	34.20	AB568417	ref. 9	
<i>S5-SLF8</i>	5.74	AB932965	this work	
<i>S5-SLF9</i>	27.11	AB932966	this work	
<i>S5-SLF10</i>	18.07	AB932967	this work	
<i>S5-SLF11</i>	11.28	AB932968	this work	
<i>S5-SLF12</i>	8.66	AB932969	this work	
<i>S5-SLF13</i>	6.59	AB932970	this work	
<i>S5-SLF15ψ</i>	undetectable	AB932971	this work	1 bp insertion at position 107
<i>S5-SLF16</i>	22.18	AB932972	this work	
<i>S5-SLF17</i>	10.36	AB932973	this work	
<i>S5-SLF18</i>	123.97	AB568423	ref. 9	synonymous with S5-FBX, identical to S10-/S22-/S22m-SLF18s
<i>S5-FBX2</i>	14.80	AB932974	this work	
total tags number	590953			

S7-haplotype

Gene	RPKM	Accession ID	References	Note
<i>S7-SLF1</i>	5.60	AB568391	ref. 9	interact with S9- and S17-RNases, identical to S19-/SC2-/S22m-SLF1s
<i>S7-SLF1B</i>	5.42	AB932975	this work	
<i>S7-SLF2</i>	3.30	AB568395	ref. 9	interact with S9-, S11-, and S19-RNases
<i>S7-SLF3</i>	2.20	AB932976	this work	
<i>S7-SLF4</i>	undetectable	AB568406	ref. 9	
<i>S7-SLF5</i>	4.29	AB568412	ref. 9	
<i>S7-SLF6</i>	4.52	AB568418	ref. 9	
<i>S7-SLF8A</i>	9.24	AB932977	this work	
<i>S7-SLF8B</i>	2.14	AB932978	this work	identical to S19-/SC2-/S22m-SLF8s
<i>S7-SLF9A</i>	25.75	AB932979	this work	interact with S19-RNase
<i>S7-SLF9Bψ</i>	10.89	AB932980	this work	deletion from position 646
<i>S7-SLF10</i>	36.14	AB932981	this work	
<i>S7-SLF11</i>	8.07	AB932982	this work	
<i>S7-SLF13</i>	27.53	AB568400	ref. 9	renamed from S7-SLF3
<i>S7-SLF14</i>	4.25	AB932983	this work	
<i>S7-SLF16</i>	25.96	AB932984	this work	
<i>S7-SLF17</i>	6.39	AB932985	this work	
<i>S7-FBX1</i>	1.26	AB932986	this work	
total tags number	634016			

Supplementary Table 1 (continued)

S9-haplotype

Gene	RPKM	Accession ID	References	Note
<i>S9-SLF1</i>	undetectable	AB568392	ref. 9	interact with S17-RNase
<i>S9-SLF3</i>	28.33	AB568401	ref. 9	
<i>S9-SLF4</i>	9.17	AB568407	ref. 9	
<i>S9-SLF5</i>	1.10	AB568413	ref. 9	
<i>S9-SLF6</i>	3.38	AB568419	ref. 9	
<i>S9-SLF7</i>	30.54	AB932987	this work	
<i>S9-SLF8</i>	5.03	AB932988	this work	
<i>S9-SLF9A</i>	2.32	AB932989	this work	
<i>S9-SLF9B</i>	7.76	AB932990	this work	
<i>S9-SLF10</i>	40.51	AB932991	this work	
<i>S9-SLF11</i>	23.05	AB932992	this work	
<i>S9-SLF12ψ</i>	undetectable	AB932993	this work	1 bp deletion at position 193
<i>S9-SLF13</i>	2.85	AB932994	this work	
<i>S9-SLF14</i>	5.33	AB932995	this work	
<i>S9-SLF15</i>	undetectable	AB932996	this work	
<i>S9-SLF16A</i>	7.85	AB932997	this work	
<i>S9-SLF16B</i>	1.33	AB932998	this work	
<i>S9-SLF17</i>	8.27	AB932999	this work	
<i>S9-FBX1</i>	16.17	AB933000	this work	
total tags number	550736			

S10-haplotype

Gene	RPKM	Accession ID	References	Note
<i>S10-SLF1</i>	—	AB933001	this work	
<i>S10-SLF3</i>	—	AB933002	this work	
<i>S10-SLF4</i>	—	AB933003	this work	
<i>S10-SLF5</i>	—	AB933004	this work	
<i>S10-SLF8</i>	—	AB933005	this work	
<i>S10-SLF10</i>	—	AB933006	this work	
<i>S10-SLF11</i>	—	AB933007	this work	
<i>S10-SLF12</i>	—	AB933008	this work	
<i>S10-SLF13</i>	—	AB933009	this work	
<i>S10-SLF14</i>	—	AB933010	this work	
<i>S10-SLF16</i>	—	AB933011	this work	
<i>S10-SLF17</i>	—	AB933012	this work	
<i>S10-SLF18</i>	—	AB933013	this work	
total tags number	no data			identical to S5-/S22-/S22m-SLF18s

Supplementary Table 1 (continued)

S11-haplotype

Gene	RPKM	Accession ID	References	Note
<i>S11-SLF1</i>	16.41	AB568393	ref. 9	interact with S17-RNase
<i>S11-SLF2</i>	1.31	AB568396	ref. 9	interact with S9-RNase
<i>S11-SLF3</i>	21.95	AB568402	ref. 9	interact with S7-RNase
<i>S11-SLF3B</i>	2.34	AB933014	this work	
<i>S11-SLF4</i>	undetectable	AB568408	ref. 9	
<i>S11-SLF5</i>	2.55	AB568414	ref. 9	
<i>S11-SLF6</i>	1.15	AB568420	ref. 9	
<i>S11-SLF7</i>	undetectable	AB933015	this work	
<i>S11-SLF8</i>	undetectable	AB933016	this work	
<i>S11-SLF9</i>	1.36	AB933017	this work	interact with S19-RNase
<i>S11-SLF10</i>	29.63	AB933018	this work	
<i>S11-SLF11</i>	8.68	AB933019	this work	
<i>S11-SLF13</i>	4.39	AB933020	this work	
<i>S11-SLF14</i>	2.57	AB933021	this work	
<i>S11-SLF15ψ</i>	2.70	AB933022	this work	26 bp deletion at position 732
<i>S11-SLF16</i>	4.94	AB933023	this work	
<i>S11-FBX1</i>	2.61	AB933024	this work	
total tags number	545880			

S17-haplotype

Gene	RPKM	Accession ID	References	Note
<i>S17-SLF1</i>	58.77	AY766153	ref. 21	synonymous with SLF-S17
<i>S17-SLF1B</i>	12.96	AY766155	ref. 21	synonymous with PaF1
<i>S17-SLF2</i>	9.23	AB568397	ref. 9	partial sequence
<i>S17-SLF3</i>	26.51	AB568403	ref. 9	
<i>S17-SLF4A</i>	4.57	AB568409	ref. 9	
<i>S17-SLF4Bψ</i>	1.42	AB933025	this work	2 bp insertion at position 802
<i>S17-SLF5</i>	2.55	AB568415	ref. 9	
<i>S17-SLF6</i>	14.98	AB568421	ref. 9	
<i>S17-SLF7ψ</i>	7.53	AB933026	this work	5 bp deletion at position 236
<i>S17-SLF8A</i>	4.27	AB933027	this work	
<i>S17-SLF8Bψ</i>	4.05	AB933028	this work	premature stop at position 535
<i>S17-SLF9A</i>	undetectable	AB933029	this work	
<i>S17-SLF9B</i>	30.29	AB933030	this work	
<i>S17-SLF10</i>	52.70	AB933031	this work	
<i>S17-SLF11</i>	28.40	AB933032	this work	
<i>S17-SLF12A</i>	undetectable	AB933033	this work	
<i>S17-SLF12B</i>	1.29	AB933034	this work	
<i>S17-SLF12Cψ</i>	2.42	AB933035	this work	943 bp insertion at position 523
<i>S17-SLF13</i>	5.07	AB933036	this work	
<i>S17-SLF14</i>	8.83	AB933037	this work	
<i>S17-SLF16</i>	23.63	AB933038	this work	
<i>S17-SLF17</i>	11.07	AB933039	this work	
total tags number	550612			

Supplementary Table 1 (continued)

S19-haplotype

Gene	RPKM	Accession ID	References	Note
<i>S19-SLF1</i>	4.38	AY766154	ref. 21	synonymous with SLF-S19, identical to S7-/SC2-/S22m-SLF1s
<i>S19-SLF1B</i>	23.67	AB933040	this work	
<i>S19-SLF2</i>	32.32	AB568398	ref. 9	
<i>S19-SLF3</i>	50.63	AB568404	ref. 9	
<i>S19-SLF4</i>	5.66	AB568410	ref. 9	
<i>S19-SLF5A</i>	15.84	AB568416	ref. 9	
<i>S19-SLF5B</i>	19.31	AB933041	this work	
<i>S19-SLF6</i>	16.77	AB568422	ref. 9	
<i>S19-SLF7ψ</i>	20.25	AB933042	this work	29 bp insertion at position 821
<i>S19-SLF8A</i>	10.82	AB933043	this work	identical to S7-/SC2-/S22m-SLF8s
<i>S19-SLF8B</i>	21.13	AB933044	this work	
<i>S19-SLF10A</i>	26.60	AB933045	this work	
<i>S19-SLF10B</i>	20.90	AB933046	this work	
<i>S19-SLF11</i>	24.07	AB933047	this work	
<i>S19-SLF12</i>	8.70	AB933048	this work	
<i>S19-SLF13</i>	14.41	AB933049	this work	
<i>S19-SLF14</i>	21.03	AB933050	this work	
<i>S19-SLF15ψ</i>	10.23	AB933051	this work	1 bp insertion at position 905
<i>S19-SLF16</i>	19.34	AB933052	this work	
<i>S19-FBX1</i>	20.96	AB933053	this work	
<i>S19-FBX2</i>	31.13	AB933054	this work	
<i>S19-FBX3</i>	6.65	AB933055	this work	
total tags number	521109			

S22-haplotype

Gene	RPKM	Accession ID	References	
<i>S22-SLF1</i>	—	AB933056	this work	
<i>S22-SLF3</i>	—	AB933057	this work	
<i>S22-SLF4</i>	—	AB933058	this work	
<i>S22-SLF5</i>	—	AB933059	this work	
<i>S22-SLF6</i>	—	AB933060	this work	
<i>S22-SLF7ψ</i>	—	AB933061	this work	31 bps deletions, frameshift
<i>S22-SLF8</i>	—	AB933062	this work	
<i>S22-SLF9</i>	—	AB933063	this work	
<i>S22-SLF10</i>	—	AB933064	this work	identical to Sm-SLF10
<i>S22-SLF11</i>	—	AB933065	this work	
<i>S22-SLF12</i>	—	AB933066	this work	
<i>S22-SLF13</i>	—	AB933067	this work	
<i>S22-SLF14</i>	—	AB933068	this work	
<i>S22-SLF15</i>	—	AB933069	this work	
<i>S22-SLF16</i>	—	AB933070	this work	
<i>S22-SLF18</i>	—	AB933071	this work	identical to S5-/S10-/S22m-/Sm-SLF18s
total tags number	no data			

Supplementary Table 1 (continued)

***S_{0m}*-haplotype**

Gene	RPKM	Accession ID	References	Note
<i>S0m-SLF1</i>	15.14	AB933072	this work	partial sequence
<i>S0m-SLF2</i>	8.15	AB933073	this work	
<i>S0m-SLF3</i>	undetectable	AB933074	this work	
<i>S0m-SLF4</i>	7.05	AB933075	this work	
<i>S0m-SLF5</i>	undetectable	AB933076	this work	1 bp insertion at position 78
<i>S0m-SLF6</i>	20.38	AB933077	this work	
<i>S0m-SLF7</i>	30.52	AB933078	this work	
<i>S0m-SLF8</i>	7.47	AB933079	this work	
<i>S0m-SLF10</i>	33.22	AB933080	this work	
<i>S0m-SLF11</i>	22.65	AB933081	this work	
<i>S0m-SLF12</i>	7.45	AB933082	this work	
<i>S0m-SLF13</i>	15.75	AB933083	this work	
<i>S0m-SLF14</i>	27.73	AB933084	this work	
<i>S0m-SLF15ψ</i>	7.26	AB933085	this work	
<i>S0m-SLF16</i>	undetectable	AB933086	this work	
<i>S0m-SLF17</i>	undetectable	AB933087	this work	
<i>S0m-FBX1</i>	60.39	AB933088	this work	
<i>S0m-FBX2</i>	undetectable	AB933089	this work	
total tags number	101431			

***S_m*-haplotype**

Gene	RPKM	Accession ID	References	Note
<i>Sm-SLF1</i>	—	AB933090	this work	identical to S5-/S10-/S22-/S22m-SLF18s
<i>Sm-SLF2</i>	—	AB933091	this work	
<i>Sm-SLF3</i>	—	AB933092	this work	
<i>Sm-SLF5</i>	—	AB933093	this work	
<i>Sm-SLF6</i>	—	AB933094	this work	
<i>Sm-SLF7</i>	—	AB933095	this work	
<i>Sm-SLF8</i>	—	AB933096	this work	
<i>Sm-SLF9A</i>	—	AB933097	this work	
<i>Sm-SLF9B</i>	—	AB933098	this work	
<i>Sm-SLF10</i>	—	AB933099	this work	
<i>Sm-SLF11</i>	—	AB933100	this work	
<i>Sm-SLF12A</i>	—	AB933101	this work	
<i>Sm-SLF12B</i>	—	AB933102	this work	
<i>Sm-SLF13</i>	—	AB933103	this work	
<i>Sm-SLF14</i>	—	AB933104	this work	
<i>Sm-SLF15</i>	—	AB933105	this work	
<i>Sm-SLF16</i>	—	AB933106	this work	
<i>Sm-SLF18</i>	—	AB933107	this work	
total tags number	no data			

Supplementary Table 1 (continued)

SC2-haplotype

Gene	RPKM	Accession ID	References	Note
<i>SC2-SLF1A</i>	—	AY766153	ref. 21	identical to S17-SLF1
<i>SC2-SLF1B</i>	—	AY766155	ref. 21	identical to S17-SLF1B
<i>SC2-SLF1C</i>	—	AY766154	ref. 21	identical to S7-/S19-/S22m-SLF1s
<i>SC2-SLF2</i>	—	AB933108	this work	identical to S17-SLF2
<i>SC2-SLF5</i>	—	AB933109	this work	1 bp (1 aa) different from S17-SLF5
<i>SC2-SLF6</i>	—	AB933110	this work	identical to S17-SLF6
<i>SC2-SLF7ψ</i>	—	AB933111	this work	identical to S17-SLF7 ψ
<i>SC2-SLF8A</i>	—	AB933112	this work	identical to S17-SLF8A
<i>SC2-SLF8Bψ</i>	—	AB933113	this work	identical to S17-SLF8B ψ
<i>SC2-SLF8C</i>	—	AB933114	this work	identical to S7-/S19-/S22m-SLF8s
<i>SC2-SLF9</i>	—	AB933115	this work	identical to S17-SLF9B
<i>SC2-SLF10</i>	—	AB933116	this work	identical to S17-SLF10
<i>SC2-SLF11</i>	—	AB933117	this work	identical to S17-SLF11
<i>SC2-SLF12</i>	—	AB933118	this work	identical to S17-SLF12B
<i>SC2-SLF13</i>	—	AB933119	this work	identical to S17-SLF13
<i>SC2-SLF14</i>	—	AB933120	this work	6 bps (5 aas) different from S17-SLF14
<i>SC2-SLF16</i>	—	AB933121	this work	identical to S17-SLF16
total tags number	no data			

S_{22m}-haplotype

Gene	RPKM	Accession ID	References	Note
<i>S22m-SLF1A</i>	—	AB933122	this work	identical to S22-SLF1
<i>S22m-SLF1B</i>	—	AB933123	this work	identical to S7-/S19-/SC2-SLF1s
<i>S22m-SLF3</i>	—	AB933124	this work	identical to S22-SLF3
<i>S22m-SLF4</i>	—	AB933125	this work	identical to S22-SLF4
<i>S22m-SLF5</i>	—	AB933126	this work	identical to S22-SLF5
<i>S22m-SLF6</i>	—	AB933127	this work	identical to S22-SLF6
<i>S22m-SLF7ψ</i>	—	AB933128	this work	identical to S22-SLF7 ψ
<i>S22m-SLF8A</i>	—	AB933129	this work	identical to S22-SLF8
<i>S22m-SLF8B</i>	—	AB933130	this work	identical to S7-/S19-/SC2-SLF8s
<i>S22m-SLF9</i>	—	AB933131	this work	identical to S22-SLF9
<i>S22m-SLF10</i>	—	AB933132	this work	identical to S0m-/S7-/S11-/S17-SLF10s
<i>S22m-SLF11</i>	—	AB933133	this work	identical to S22-SLF11
<i>S22m-SLF12</i>	—	AB933134	this work	identical to S22-SLF12
<i>S22m-SLF13</i>	—	AB933135	this work	identical to S22-SLF13
<i>S22m-SLF14</i>	—	AB933136	this work	identical to S22-SLF14
<i>S22m-SLF16</i>	—	AB933137	this work	identical to S22-SLF16
<i>S22m-SLF17</i>	—	AB933138	this work	identical to S0m-/S7-/S9-/S10-/S17-SLF17s
<i>S22m-SLF18</i>	—	AB933139	this work	identical to S5-/S10-/S22-/Sm-SLF18s
total tags number	no data			

ψ following a gene name indicates pseudogene. For *SLFs* of the *S₅*-, *S₇*-, *S₉*-, *S₁₁*-, *S₁₇*-, *S₁₉*-, and *S_{0m}*-haplotypes, relative expression levels are shown in RPKM (Reads Per Kilobase of exon model per Million mapped reads), and total numbers of ESTs are shown at the bottom of each table. GenBank accession IDs and references are also shown.

Supplementary Table 2 List of primer sequences

	Name	Primer	Target / Purpose
Type-7 specific amplification	SLF7-F1	CAA AGT ATA TTT TAC ATT GCA TTG CAG A	F Type-7 <i>SLFs</i> / genomic PCR
	SLF7-F2	GGG TTT TAG AAG ACT ATG GCG ATC	F Type-7 <i>SLFs</i> / 3' RACE, genomic PCR
	SLF7-F3	GTT GAG AAA GTG GAA GAT CGA TGT	F Type-7 <i>SLFs</i> / 3' RACE, genomic PCR
	SLF7-F4	ATG ACT CTC TAA CAC TGA TCT GTT ACC G	F Type-7 <i>SLFs</i> / 3' RACE, expression analysis
	SLF7-R1	GAG AAG GGT AAG CTC TAT TTA AAT TGG	R Type-7 <i>SLFs</i> / genomic PCR
	SLF7-R2	GTA CTC CTT CAT TGT CCA AAT TTC CA	R Type-7 <i>SLFs</i> / 5' RACE, genomic PCR
	SLF7-R3	TTC CGT TCT TCC AAA CCA ATG AG	R Type-7 <i>SLFs</i> / 5' RACE, genomic PCR
	SLF7-R4	TGA GCT CTC TAA AAA TTT TGA ACT TGA GTA CTA	R Type-7 <i>SLFs</i> / genomic PCR, expression analysis
Type-8 specific amplification	SLF8-F1	TCA AAG GAT CGG AGG TAT TAC AGA GA	F Type-8 <i>SLFs</i> / genomic PCR
	SLF8-F2	GAA ACA TTT TTT ATT GCA TTG AAG AGA AC	F Type-8 <i>SLFs</i> / genomic PCR
	SLF8-F3	GGT TTT GAC TCG ATT GTG AAG GAC T	F Type-8 <i>SLFs</i> / 3' RACE
	SLF8-F4	AAG CAA AAG TGG ACT TCT TGT TTC CTA C	F Type-8 <i>SLFs</i> / 3' RACE, expression analysis
	SLF8-R1	ATT TAG GCT GAA AAC TAC TTT CCA ATC A	R Type-8 <i>SLFs</i> / genomic PCR, expression analysis
	SLF8-R2	CCT TTT TTG GAA TTG AAA TCA AGC	R Type-8 <i>SLFs</i> / 5' RACE
	SLF8-R3	CTC CAA GAA TCA AAA CGC AAA TCA	R Type-8 <i>SLFs</i> / 5' RACE
Type-9 specific amplification	SLF9-F1	ATC CAA CTG AAA TGA GGT CTT AAA GCA	F Type-9 <i>SLFs</i> , <i>S19-FBX1</i> , 2 and <i>S0m-FBX1</i> / genomic PCR
	SLF9-F2	GCC GTG GTA CCA AAA AGG CAA	F Type-9 <i>SLFs</i> , <i>S0m-/S19-FBX1</i> and 2 / genomic PCR
	SLF9-F3	GAA GTT AAG GAA TTA GAT TTG CAT GGT CT	F Type-9 <i>SLFs</i> / 3' RACE, expression analysis
	SLF9/10-F1	AAT GAA GGA ATA CGG TGA AAA GGA G	F Type-9 and -10 <i>SLFs</i> , <i>S0m-/S19-FBX1</i> and 2 / 3' RACE
	SLF9/10-F2	GAA TCC CCA TTA GCA GTT TGG AAG	F Type-9 and -10 <i>SLFs</i> , <i>S0m-/S19-FBX1</i> and 2 / 3' RACE
	SLF9-R1	CCA TTG ATC TGT AAG TTG TTG CTG C	R Type-9 <i>SLFs</i> / genomic PCR
	SLF9-R2	GAA GTT CTA TGC AAT CCT TAT TTC TTG G	R Type-9 <i>SLFs</i> and <i>S0m-/S19-FBX1</i> / 5' RACE
	SLF9-R3	CGA ATT AAG ATC ATA GGC AAT CAA ATG T	R Type-9 <i>SLFs</i> and <i>S0m-/S19-FBX1</i> / 5' RACE
	SLF9-R4	AAT ATG GCT TCC ATT GAT CTG TAA GTT G	R Type-9 <i>SLFs</i> / genomic PCR, expression analysis
Type-10 specific amplification	SLF10-F1	CCG AAG TAA GAT CCA ACT AAA ATG ATG T	F Type-10 <i>SLFs</i> / genomic PCR
	SLF10-F2	GAA GTC CAG GAA TTA GAT TTA CAT GGT TA	F Type-10 <i>SLFs</i> / 5' RACE, expression analysis
	SLF9/10-F1	AAT GAA GGA ATA CGG TGA AAA GGA G	F Type-9 and -10 <i>SLFs</i> , <i>S0m-/S19-FBX1</i> and 2 / 3' RACE
	SLF9/10-F2	GAA TCC CCA TTA GCA GTT TGG AAG	F Type-9 and -10 <i>SLFs</i> , <i>S0m-/S19-FBX1</i> and 2 / 3' RACE
	SLF10-R1	TAG TAT TCA GAT GTG TTA GGT TGT TGC C	R Type-10 <i>SLFs</i> / genomic PCR, expression analysis
	SLF10-R2	ATT AAA TAC GAT AGT GGT AAG GGA ATC TGT	R Type-10 <i>SLFs</i> / 5' RACE
	SLF10-R3	CAG CAG TGG TGG TCA AAT GTG G	R Type-10 <i>SLFs</i> / 5' RACE
Type-11 specific amplification	SLF11-F1	ATG TTT GAC GGG TCG CTC CTT	F Type-11 <i>SLFs</i> / genomic PCR
	SLF11-F2	ATT ACT ACA CAA TCA TTC CCA TAT AGC ACA	F Type-11 <i>SLFs</i> / 3' RACE, genomic PCR
	SLF11-F3	TTG TTG GAG AGA CCT GGA TCT TAC G	F Type-11 <i>SLFs</i> / 3' RACE, expression analysis
	SLF11-R1	AAT TAA AAG TGT GGA AAT CTT CAC AAA GAA	R Type-11 <i>SLFs</i> / 5' RACE, genomic PCR
	SLF11-R2	GAG GCC ATA ACC TTT ATT GTC CCA	R Type-11 <i>SLFs</i> / 5' RACE, genomic PCR, expression analysis
Type-12 specific amplification	SLF12-F1	AAG GGA GTA AAG CTG CTA TTT TGA	F Type-12 <i>SLFs</i> / genomic PCR
	SLF12-F2	CTT CCA GTA AAG TTT CTC TTG CGA TTC	F Type-12 <i>SLFs</i> / 3' RACE, expression analysis
	SLF12-F3	TCA CGC TCT CAC AAA ATC TTC CAC T	F Type-12 <i>SLFs</i> / 3' RACE
	SLF12-R1	AGT ATT AAA ATC CCA CGA TGA AAA AG	R Type-12 <i>SLFs</i> / genomic PCR, expression analysis
	SLF12-R2	TAA TTG TGT ATT TCT TTG TCC AAG ACT CAT	R Type-12 and -4 <i>SLFs</i> / 5' RACE
	SLF12-R3	GGA ACT TAC ATT CAT CAC AGC AAC CAG	R Type-12 <i>SLFs</i> / 5' RACE
Type-13 specific amplification	SLF13-F1	AAA TGT TAT TAC ACA CAC AAT CCT TGA A	F Type-13 <i>SLFs</i> / 3' RACE, genomic PCR
	SLF13-F2	TGG TTT GAT TCT TTT GAC AGA CTC GTT	F Type-13 <i>SLFs</i> / 3' RACE, expression analysis
	S7-SLF13-R	CTT AAG TTG GGG TGT TTT ACC TTC TC	R Type-13 <i>SLFs</i> / genomic PCR
	SLF13-R1	AAG ATC TCC ATA CAC ACG TGA AAC TCT AAT	R Type-13 <i>SLFs</i> / 5' RACE
	SLF13-R3	AAA ACA GAG AAG TAA ACG CAT TTG TCC	R Type-13 <i>SLFs</i> / 5' RACE, genomic PCR, expression analysis
Type-14 specific amplification	SLF14-F1	AGC TGA TAC TAC TCA AAG AAA CCT AAC TC	F Type-14 <i>SLFs</i> / genomic PCR
	SLF14-F2	ATT CTC TTC AAG CGC TCG TTC AA	F Type-14 <i>SLFs</i> / 3' RACE
	SLF14-F3	ATC TCA TTG GTT CGG AAA CAC AAA TAC	F Type-14 <i>SLFs</i> / 3' RACE, expression analysis
	SLF14/16-F1	TGT CTC TCC AGA TTT AGA AGT GCC ATA	F Type-14 and -16 <i>SLFs</i> / 3' RACE
	SLF14/16-F2	GGA GAG AGG TAG AAA ATG TGG ATC AAC	F Type-14 and -16 <i>SLFs</i> / 3' RACE
	SLF14-R1	TTA TTA TTG ATA GGC TTT TAC ACA ATT TAA	R Type-14 <i>SLFs</i> / genomic PCR
	SLF14-R2	AAG CAT TTC TAA TAA CTG AAC TAA CTT AGG GA	R Type-14 <i>SLFs</i> / 5' RACE, genomic PCR, expression analysis
	SLF14-R3	GAA CGA TGG AAA CCT AGT GGA CTG	R Type-14 <i>SLFs</i> / 5' RACE
	SLF14-R4	GTT GCT GGA TTA GAT AAT ACT GTA GTT ACT TT	R Type-14 <i>SLFs</i> / 5' RACE

	SLF14/16-R1	ACT ACG AGG CCA TAA CAC TTC CTG T	R	Type-14 and -16 <i>SLFs</i> / 5' RACE
	SLF14/16-R2	CTG GAG AAA CAT GGT GAA GGT CAT AA	R	Type-14 and -16 <i>SLFs</i> / 5' RACE
Type-15 specific amplification	SLF15-F1	TTA GAG CTA TTA CAC TGC AAA GGA ACC A	F	Type-15 <i>SLFs</i> / genomic PCR
	SLF15-F2	CTG AAG TAC AAT GTG CAT TGC TTT GA	F	Type-15 <i>SLFs</i> / 3' RACE
	SLF15-F3	GTG AAA TTA TGG AAA GAT TGC CTA AAG A	F	Type-15 <i>SLFs</i> / 3' RACE
	SLF15-F4	GGA TAA TGA TGG AAT ACG GTG TGG A	F	Type-15 <i>SLFs</i> / 3' RACE, expression analysis
	SLF15-R1	GGA ATG CAT AAC CTG AAA CGA CAA	R	Type-15 <i>SLFs</i> / 5' RACE, genomic PCR
	SLF15-R2	TTT TGG GCA TTT GGT ACT ATG CTC T	R	Type-15 <i>SLFs</i> / 5' RACE
	SLF15-R3	CCC AGT TAA AAA GTT ATC GAG AAG TAG GT	R	Type-15 <i>SLFs</i> / 5' RACE, genomic PCR, expression analysis
Type-16 specific amplification	SLF16-F1	ATG TCC CTA AAA TTC TAC AGT GAA CAT TGA	F	Type-16 <i>SLFs</i> / genomic PCR
	SLF16-F2	GTG CAT TGG TAT CCG TGT GGC	F	Type-16 <i>SLFs</i> / 3' RACE, expression analysis
	SLF14/16-F1	TGT CTC TCC AGA TTT AGA AGT GCC ATA	F	Type-16 and -14 <i>SLFs</i> / 3' RACE
	SLF14/16-F2	GGA GAG AGG TAG AAA ATG TGG ATC AAC	F	Type-16 and -14 <i>SLFs</i> / 3' RACE
	SLF16-R1	TAG CTA AAG AAT ATG AAT GAA AAG AAA GAT CC	R	Type-16 <i>SLFs</i> / genomic PCR
	SLF16-R2	CCG AAG AGA GAG TTA TAG AAA TGC TGG T	R	Type-16 <i>SLFs</i> / genomic PCR
	SLF16-R3	ATG CTG GTT TTA TAT TTT GAA GTT AGA GAG	R	Type-16 <i>SLFs</i> / 5' RACE, genomic PCR, expression analysis
	SLF14/16-R1	ACT ACG AGG CCA TAA CAC TTC CTG T	R	Type-16 and -14 <i>SLFs</i> / 5' RACE
	SLF14/16-R2	CTG GAG AAA CAT GGT GAA GGT CAT AA	R	Type-16 and -14 <i>SLFs</i> / 5' RACE
Type-17 specific amplification	SLF17-F1	GCG CAC ACA TAA AAA ACA TTT CCT AA	F	Type-17 <i>SLFs</i> / genomic PCR
	SLF17-F2	TAT GTG CAT TGG TAT CCC AGT GCT	F	Type-17 <i>SLFs</i> / 3' RACE, expression analysis
	SLF17-R1	TAA GTC AAG TCT CTA CCA GTT GGA CTC G	R	Type-17 <i>SLFs</i> / genomic PCR, expression analysis
	SLF17-R2	TAT GCT CGC GGT CTT TTG GAA	R	Type-17 and -16 <i>SLFs</i> / 5' RACE
	SLF17-R3	AAC CAA TGA GAG GCC CCT TTG TA	R	Type-17 <i>SLFs</i> / 5' RACE
Type-18 specific amplification	S5-FBX-F1	CCT CAT GCA ATG AAT TGC TCA AGC AGT	F	Type-18 <i>SLFs</i> / genomic PCR
	S5-FBX-F2	GAT CAA TCG ACT GCT CAA GGA AAA	F	Type-18 <i>SLFs</i> / 3' RACE
	S5-FBX-SF	TCG TCG GTT GTT TGG ATA ATG AAA G	F	Type-18 <i>SLFs</i> / 3' RACE, expression analysis
	S5-FBX-R1	TTG ATT GGA CTC TAT TGG AAT TAG GTG	R	Type-18 <i>SLFs</i> / genomic PCR
	S5-FBX-R2	TCA AAA ACC AAC AAA TAA CCC TTC CAA	R	Type-18 <i>SLFs</i> / 5' RACE
	S5-FBX-R3	CAA TCC AAT GGC AAG CTC CCT TGT	R	Type-18 <i>SLFs</i> / 5' RACE
	S5-FBX-SR	TAC ATG AGA GTT ACA AAT CAA CCC AAA G	R	Type-18 <i>SLFs</i> / genomic PCR, expression analysis
Allele-specific amplification for SLF	S5-SLF9-F	ACA ATA TAG GAA TGC CAG ATG CT	F	<i>S5-SLF9</i> / genotyping
	S5-SLF9-R	CTC TTC ACG ACA GAA AGT TTT TCA	R	<i>S5-SLF9</i> / genotyping
	S5-SLF12-F	CCG ATT ATG AAA TTG CAT GAA GAT ATT G	F	<i>S5-SLF12</i> / genotyping
	S5-SLF12-R	CCG TAC AGT TAA GAT CAT CAC CAT TAC C	R	<i>S5-SLF12</i> / genotyping
	S5-SLF16-F	AAG GAG TAT GGT ATA AAC GAG TCT TGG TG	F	<i>S5-SLF16</i> / genotyping
	S5-SLF16-R	AAG GAG TAT GGT ATA AAC GAG TCT TGG TG	R	<i>S5-SLF16</i> / genotyping
	S5-FBX-SF	CAC CTA ATT CCA ATA GAG TCC AAT CAA TA	F	<i>S5-, S10-, S22-, Sm-SLF18</i> / genotyping, expression analysis
	S5-FBX-SR	TAC ATG AGA GTT ACA AAT CAA CCC AAA G	R	<i>S5-, S10-, S22-, Sm-SLF18</i> / genotyping, expression analysis
	S5-FBX2-F1	AAA CCG AAA CAA TAT TCC CCA AAG TAG	F	<i>S5-FBX2</i> / genomic cloning
	S5-FBX2-R1	AAA AGG TTG CAG AAA TCC TGG TGA TA	R	<i>S5-FBX2</i> / genomic cloning
	S5-FBX2-F2	ACG GCC TGG TAA TCC TGA ATG AG	F	<i>S5-FBX2</i> / genotyping, expression analysis
	S5-FBX2-R2	GAT AAT CCT CTC CAA AGT TGT TGA ACG	R	<i>S5-FBX2</i> / genotyping, expression analysis
	S7-SLF11-F	GAT GAA CTC AAG GGA TTC AAT TTC CA	F	<i>S7-SLF11</i> / genotyping
	S7/S11-SLF11-R	TTC TCA AAG AAA TTG TAA CAG ATT AGT GCC	R	<i>S7-SLF11</i> / genotyping
	S7-SLF13-F	CCA AGG AGG GAG AGT AGT CCA ATG	F	<i>S7-SLF13</i> / genotyping
	S7-SLF13-R	CTT AAG TTG GGG TGT TTT ACC TTC TC	R	<i>S7-SLF13</i> / genotyping
	S7-SLF14-F	TTT GCT ATT GAA TCA CCA TTG GCT ATC	F	<i>S7-SLF14</i> / genotyping
	S7-SLF14-R	TGC GCG AGT ATT ATT GAG CTT GTG	R	<i>S7-SLF14</i> / genotyping
	S7-SLF16-F	GTT AGT CAA CAT TGC ATC ACA AGA AAG A	F	<i>S7-SLF16</i> / genotyping
	S7-SLF16-R	TTC CTT CGC AAT CTG TTA AAA CCA A	R	<i>S7-SLF16</i> / genotyping
	S7-FBX1-F1	GGT TTG ATT GCT TTG ACA GAT TTT GAA	F	<i>S7-FBX1</i> / 3' RACE, genotyping, expression analysis
	S7-FBX1-R1	AAG AAT TAC CAT TGT GTC TGT GTC TGT ATC	R	<i>S7-FBX1</i> / 5' RACE, genotyping, expression analysis
	S7-FBX1-F2	ATG GAT TCA GAT TGG ATT CCC GT	F	<i>S7-FBX1</i> / 3' RACE, genomic PCR
	S7-FBX1-R2	TCC CAT GTG AAA GAT CTC GAA ATG T	R	<i>S7-FBX1</i> / 5' RACE, genomic PCR
	S9-SLF7-F	AAT CGA TCC TAC AAA AGA CTT CAT GGA G	F	<i>S9-SLF7</i> / genotyping
	S9-SLF7-R	ACA AAA AGA TGT TCA AAA GCA GGT GTT	R	<i>S9-SLF7</i> / genotyping
	S9-SLF10-F	AAA TGG GCA ATG CAC AAT ACA AGT T	F	<i>S9-SLF10</i> / genotyping
	S9-SLF10-R	CCA GGC CAT AAG ACT TTC CAT CC	R	<i>S9-SLF10</i> / genotyping

	S9-SLF11-F	CTA AAA CAT GGT ACG CGC TCG TC	F <i>S9-SLF11</i> / genotyping
	S9-SLF11-R	AAC TAA CTT GTA ATA ATT TTC AGC CAG GG	R <i>S9-SLF11</i> / genotyping
	S9-SLF13-F	TTT ACG GTG TTG GAT TTG GCG	F <i>S9-SLF13</i> / genotyping
	S9-SLF13-R	ATA TTG CCA AAG GGG ATT CGA C	R <i>S9-SLF13</i> / genotyping
	S9-SLF15-F	GTG GTC TTG GTG ATG ATG ATC TTT G	F <i>S9-SLF15</i> / genotyping
	S9-SLF15-R	AAA AAA AAG AAT TAC CAT TGT ATT TGC AC	R <i>S9-SLF15</i> / genotyping
	S9-FBX1-F1	AGA GTG CAA CAT CGT ATA ACA GAG CC	F <i>S9-FBX1</i> / 3' RACE, genomic PCR
	S9-FBX1-F2	CAA TGG ACC TAG TTA TGG CCT CCT	F <i>S9-FBX1</i> / 3' RACE, genotyping, expression analysis
	S9-FBX1-R1	AGA TTT GGT GGG TAT TCG GTG CTA	R <i>S9-FBX1</i> / genomic PCR, genotyping, expression analysis
	S11-SLF8-F	CAG CGA CGA TCT TAT CCC CAT TA	F <i>S11-SLF8</i> / genotyping
	S11-SLF8-R	CAT CAT TAT CCA AAT GTC CAT TAA ATC A	R <i>S11-SLF8</i> / genotyping
	S11-SLF11-F	CTG ATG AAC TCA AGG AAT TCA ATT TCA G	F <i>S11-SLF11</i> / genotyping
	S7/S11-SLF11-R	TTC TCA AAG AAA TTG TAA CAG ATT AGT GCC	R <i>S11-SLF11</i> / genotyping
	S11-FBX1-F1	TTA CCC CTA CCC AGA GCC TGT G	F <i>S11-FBX1</i> / 3' RACE, genomic PCR
	S11-FBX1-F2	GGA TTA AAA AGT ACA CAA TTA CAG GCC	F <i>S11-FBX1</i> / 3' RACE, genotyping, expression analysis
	S11-FBX1-R1	CGT TAT CTT ACA AGC CAA ACA AGC	R <i>S11-FBX1</i> / 5' RACE, genotyping, expression analysis
	S11-FBX1-R2	GGA TAC CCG TCG TTA ACA ATG	R <i>S11-FBX1</i> / 5' RACE, genomic PCR
	SLF8-F2	GAA ACA TTT TTT ATT GCA TTG AAG AGA AC	F <i>S17-SLF8A</i> / genotyping
	S17-SLF8A-F	GAA TCT TTT GTT GGA TCC ATC TTG TTA	R <i>S17-SLF8A</i> / genotyping
	S17-SLF12B-F	AAG ATG CTG GAC GGA ATT ATT ATG AAG	F <i>S17-SLF12B/Cψ</i> / genotyping
	S17-SLF12B-R	CTC TCT CCC TAT TCA GGA TAC CCG A	R <i>S17-SLF12B/Cψ</i> / genotyping
	S19-SLF10B-F	CAA GTA GTC CAA CAG AAA AAT TCA CG	F <i>S19-SLF10B</i> / genotyping
	S19-SLF10B-R	TTT CAT AAT GGA GGC AAA AGA AGC	R <i>S19-SLF10B</i> / genotyping
	S19-FBX1-F	CCT GGT GAT TTT GTG TAA GTC GCT	F <i>S19-FBX1</i> , <i>S0m-FBX1</i> / 3' RACE, genotyping
	S19-FBX1-R	GTT AGG TTG TTG CCG AAA TCA AAG	R <i>S19-FBX1</i> / genomic PCR, genotyping
	S19-FBX2-F	CTT CTT CCT ATT GAA TCC CCG TTG	F <i>S19-FBX2</i> , <i>S0m-FBX2</i> / 3' RACE, genomic PCR, genotyping
	S19-FBX2-R	GTG AGT ATC TAA AAT GGT TGA ACT TCA GC	R <i>S19-FBX2</i> / genomic PCR, genotyping
	S19-FBX3-F1	AAA TAC ACA CTA GAA TGT TAA GAG TCA	F <i>S19-FBX3</i> / 3' RACE, genomic PCR
	S19-FBX3-F2	AGA AGT TTA CTG GAT AGA TCA TGG	F <i>S19-FBX3</i> / 3' RACE, genomic PCR, genotyping
	S19-FBX3-R1	GCT AAT GGA GAT ACA ATA CGA AGA	R <i>S19-FBX3</i> / genomic PCR, genotyping
	SLF9-F2	GCC GTG GTA CCA AAA AGG CAA	F <i>S0m-FBX2</i> , <i>Type-9 SLFs</i> , <i>S19-FBX1</i> and <i>2</i> / genomic PCR
	S0m-FBX2-R	TGG AAG GAA ATG GAG ATA ATG TTG AAA	R <i>S0m-FBX2</i> / genomic PCR
<i>S-RNases</i> amplification	SRNC1-U	HWK GCA ACT MGT ITT AAC ATG GCC	F <i>S-RNase</i> conserved region 1
	SRNC5-L	WSG VKY RAA RCA TAT ICC TAT CTC	R <i>S-RNase</i> conserved region 5
Allele-specific amplification for <i>S-RNase</i>	S0m-RN-F	ACT TCA GAA TGT TTC AGT TTC AG	F <i>S0m-RNase</i> / genotyping
	S0m-RN-R	AAC AGG TTT ACT TTT GCT AAT TAT	R <i>S0m-RNase</i> / genotyping
	S22-RN-F	TTT TCG TCT AGA GTT CTG TCC TGG C	F <i>S22-RNase</i> / genotyping
	S22-RN-R	CGC ATT TGA GGT CAG GAT CTT TG	R <i>S22-RNase</i> / genotyping
	S10-RN-F	GGC CAG ATA ATG AGC AAA GAC GT	F <i>S10-RNase</i> / genotyping
	S10-RN-R	CAA AAA AGA AAG AAA TGG AAA GTC AAA C	R <i>S10-RNase</i> / genotyping
	Sm-RN-F	AGA ATT GCC CGA TAA AAC CG	F <i>Sm-RNase</i> / genotyping
	Sm-RN-R	TTC TGA TTG ACT GTA GAC ATC TG	R <i>Sm-RNase</i> / genotyping
Vector construction	S5-T3-BamHI-F	AAG GAT CCA TGA CGG CCA TGA AGA AAT TGC	F Cloning of <i>S5-SLF3</i> ORF
	S11-T3-SacI-R	AAT GAG CTC TAA AAA TTC TGA ACT TGT GTA CTA C	R Cloning of <i>S5-SLF3</i> ORF
	T9-BamHI-F	TGG ATC CAT GAA GGA ATT GCC CCA AG	F Cloning of <i>S7-</i> and <i>S11-SLF9s</i> ORF
	T9-SacI-R	AGA GCT CTC CCC ACT TCT AAA ATT GTT CAA	R Cloning of <i>S7-</i> and <i>S11-SLF9s</i> ORF
Transgene detection	SLF3-F1	CAT TCA ATT GTC CTA GTG TGA TGG AGT	F <i>S5-SLF3</i> -transgene / genotyping, expression analysis
	SLF9/10-F1	AAT GAA GGA ATA CGG TGA AAA GGA G	F <i>S7-</i> and <i>S11-SLF9</i> -transgene / genotyping, expression analysis
	NosR	ACC GGC AAC AGG ATT CAA TC	R <i>Nos</i> terminator / genotyping, expression analysis
RT-PCR control	UB-F	CCT AAC CGG CAA AAC CAT CAC CT	F <i>Petunia ubiquitin</i> / expression analysis
	UB-R	GCA CTT ATC AAC AAC AGG ACG ACA ACA	R <i>Petunia ubiquitin</i> / expression analysis

Supplementary Table 3 Summary of pairwise peptide-sequence identities among S-genes

(a) Type-1 SLFs																			
Max. = 100.0 % Min. = 86.1 % Ave. = 92.0 %																			
	S9-SLF1	Sm-SLF1	S22-SLF1	PIS11-SLF1	S11-SLF1	S7-SLF1	S19-SLF1	S10-SLF1	S5-SLF1	S0m-SLF1	S19-SLF1B	PIS2-SLF1	S7-SLF1B	PIS13-SLF1	PIS7-SLF1	PIS1-SLF1	S17-SLF1B	S17-SLF1	PIS3-SLF1
PIS5-SLF1	97.2 %	96.1 %	90.2 %	92.8 %	92.5 %	97.2 %	97.2 %	98.2 %	97.7 %	97.9 %	97.9 %	94.9 %	94.3 %	90.0 %	90.0 %	89.5 %	89.7 %	86.6 %	86.4 %
S9-SLF1	-	96.4 %	90.2 %	92.5 %	92.5 %	97.4 %	97.4 %	97.9 %	97.4 %	97.9 %	97.7 %	94.6 %	92.8 %	90.2 %	89.7 %	89.7 %	89.2 %	86.3 %	86.4 %
Sm-SLF1		-	90.5 %	92.0 %	91.8 %	99.0 %	99.0 %	97.9 %	96.9 %	96.9 %	96.7 %	94.6 %	94.1 %	90.2 %	89.5 %	89.2 %	88.9 %	86.6 %	86.1 %
S22-SLF1			-	91.2 %	91.0 %	91.0 %	91.0 %	90.5 %	91.0 %	91.3 %	90.7 %	92.8 %	93.0 %	91.0 %	88.9 %	88.2 %	88.4 %	87.6 %	86.6 %
PIS11-SLF1				-	97.4 %	93.1 %	93.1 %	92.8 %	93.3 %	93.1 %	93.1 %	93.3 %	96.1 %	91.0 %	89.9 %	88.9 %	89.5 %	87.6 %	87.4 %
S11-SLF1					-	92.8 %	92.8 %	92.8 %	93.1 %	92.8 %	93.1 %	93.3 %	95.6 %	91.8 %	90.7 %	89.2 %	89.2 %	87.3 %	87.1 %
S7-SLF1						-	100.0 %	99.0 %	97.9 %	97.9 %	97.7 %	95.6 %	93.1 %	91.3 %	90.5 %	90.2 %	90.0 %	87.4 %	87.1 %
S19-SLF1							-	99.0 %	97.9 %	97.9 %	97.7 %	95.6 %	93.1 %	91.3 %	90.5 %	90.2 %	90.0 %	87.4 %	87.1 %
S10-SLF1								-	98.5 %	97.9 %	98.7 %	95.1 %	93.1 %	90.7 %	90.5 %	90.2 %	90.0 %	86.9 %	86.6 %
S5-SLF1									-	97.9 %	98.2 %	95.6 %	93.3 %	91.3 %	90.2 %	90.2 %	89.7 %	87.4 %	87.4 %
S0m-SLF1										-	97.7 %	95.6 %	95.1 %	90.7 %	90.5 %	89.7 %	90.2 %	87.4 %	87.7 %
S19-SLF1B											-	94.9 %	93.3 %	91.5 %	90.7 %	90.7 %	90.5 %	87.6 %	87.4 %
PIS2-SLF1												-	95.4 %	92.3 %	90.2 %	89.7 %	90.5 %	88.7 %	88.4 %
S7-SLF1B													-	92.8 %	91.5 %	90.5 %	90.2 %	87.9 %	88.7 %
PIS13-SLF1														-	92.0 %	91.0 %	90.5 %	88.7 %	89.5 %
PIS7-SLF1															-	91.5 %	91.0 %	87.1 %	88.7 %
PIS1-SLF1																-	89.5 %	87.4 %	88.4 %
S17-SLF1B																	-	88.1 %	87.4 %
S17-SLF1																		-	86.4 %

(b) Type-2 SLFs							
Max. = 88.3 % Min. = 72.8 % Ave. = 82.4 %							
	S17-SLF2	S11-SLF2	S5-SLF2	S19-SLF2	Sm-SLF2	S0m-SLF2	PIS1-SLF2
S7-SLF2	87.1 %	84.7 %	85.2 %	83.9 %	83.5 %	85.9 %	72.8 %
S17-SLF2	-	84.4 %	85.3 %	85.3 %	83.8 %	86.2 %	74.5 %
S11-SLF2		-	83.3 %	82.6 %	82.8 %	86.2 %	73.5 %
S5-SLF2			-	85.1 %	84.6 %	83.7 %	74.7 %
S19-SLF2				-	86.2 %	87.7 %	75.0 %
Sm-SLF2					-	88.3 %	75.2 %
S0m-SLF2						-	76.0 %

(c) Type-3 SLFs											
Max. = 99.0 % Min. = 72.0 % Ave. = 88.9 %											
	S17-SLF3	S10-SLF3	S22-SLF3	S19-SLF3	PIS2-SLF3	S0m-SLF3	Sm-SLF3	S9-SLF3	S11-SLF3	S11-SLF3B	S7-SLF3
S5-SLF3	99.0 %	99.0 %	98.7 %	98.2 %	97.9 %	97.1 %	97.1 %	97.4 %	89.0 %	76.2 %	73.4 %
S17-SLF3	-	97.9 %	97.7 %	97.1 %	96.9 %	96.1 %	96.1 %	96.4 %	88.7 %	76.5 %	73.6 %
S10-SLF3		-	98.2 %	97.7 %	97.9 %	97.1 %	96.6 %	97.4 %	89.5 %	76.5 %	73.6 %
S22-SLF3			-	98.4 %	97.9 %	96.9 %	95.8 %	96.1 %	89.0 %	76.0 %	73.6 %
S19-SLF3				-	97.1 %	96.4 %	95.6 %	95.6 %	88.2 %	76.0 %	73.6 %
PIS2-SLF3					-	97.7 %	96.6 %	95.8 %	89.5 %	76.2 %	73.4 %
S0m-SLF3						-	95.8 %	95.6 %	88.2 %	76.5 %	73.6 %
Sm-SLF3							-	95.1 %	87.4 %	74.7 %	72.0 %
S9-SLF3								-	89.3 %	75.7 %	72.6 %
S11-SLF3									-	74.4 %	72.3 %
S11-SLF3B										-	85.1 %

(d) Type-4 SLFs									
Max. = 97.3 % Min. = 89.4 % Ave. = 93.4 %									
	S11-SLF4	S0m-SLF4	S5-SLF4	PIS2-SLF4	S7-SLF4	S17-SLF4Bψ	S9-SLF4	S17-SLF4	S10-SLF4
S22-SLF4	93.7 %	91.9 %	94.9 %	93.7 %	95.2 %	93.9 %	94.2 %	92.9 %	89.4 %
S11-SLF4	-	91.8 %	94.8 %	94.0 %	95.8 %	94.5 %	94.0 %	93.3 %	89.6 %
S0m-SLF4		-	93.3 %	92.6 %	93.5 %	92.6 %	93.3 %	92.1 %	89.9 %
S5-SLF4			-	96.3 %	95.3 %	95.3 %	95.5 %	94.5 %	90.9 %
PIS2-SLF4				-	95.8 %	94.5 %	94.8 %	93.8 %	90.6 %
S7-SLF4					-	97.3 %	95.8 %	94.8 %	91.6 %
S17-SLF4Bψ						-	94.8 %	94.0 %	90.4 %
S9-SLF4							-	93.8 %	90.9 %
S17-SLF4								-	92.3 %
S10-SLF4									-

100%
more than 90 %, less than 100 %
more than 80 %, less than 90 %
more than 70 %, less than 80 %
less than 70 %

Supplementary Table 3 (Continued)

(e) Type-5 SLFs Max. = 99.2 % Min. = 91.2 % Ave. = 95.5 %

	PIS3-SLF5	S11-SLF5	S0m-SLF5	S17-SLF5	PIS2-SLF5	S5-SLF5	S19-SLF5	S9-SLF5	Sm-SLF5	S10-SLF5	S22-SLF5	S5-SLF5B	S19-SLF5B
S7-SLF5	99.2 %	98.2 %	98.5 %	98.2 %	98.2 %	98.2 %	97.9 %	97.2 %	96.4 %	95.1 %	94.3 %	92.5 %	93.0 %
PIS3-SLF5	-	97.9 %	98.2 %	97.9 %	97.9 %	97.9 %	97.7 %	96.9 %	96.1 %	94.8 %	94.3 %	92.3 %	92.7 %
S11-SLF5		-	98.2 %	97.9 %	97.9 %	97.9 %	97.7 %	96.9 %	96.1 %	94.8 %	94.3 %	92.5 %	93.0 %
S0m-SLF5			-	99.2 %	98.2 %	98.2 %	97.9 %	97.2 %	96.6 %	95.4 %	94.6 %	92.8 %	93.3 %
S17-SLF5				-	97.9 %	97.9 %	97.7 %	96.9 %	96.1 %	95.1 %	94.1 %	92.3 %	92.7 %
PIS2-SLF5					-	98.2 %	97.9 %	96.9 %	96.6 %	94.8 %	94.6 %	92.3 %	92.7 %
S5-SLF5						-	97.7 %	96.9 %	96.1 %	94.8 %	94.1 %	92.5 %	93.0 %
S19-SLF5							-	96.6 %	95.9 %	94.6 %	93.8 %	92.0 %	92.5 %
S9-SLF5								-	95.6 %	94.3 %	93.0 %	91.8 %	92.2 %
Sm-SLF5									-	95.4 %	95.4 %	92.8 %	93.3 %
S10-SLF5										-	93.6 %	92.8 %	93.3 %
S22-SLF5											-	91.2 %	91.7 %
S5-SLF5B												-	99.0 %

100%
more than 90 %, less than 100 %
more than 80 %, less than 90 %
more than 70 %, less than 80 %
less than 70 %

(f) Type-6 SLFs Max. = 99.0 % Min. = 93.1 % Ave. = 94.9 %

	S0m-SLF6	Sm-SLF6	S7-SLF6	S11-SLF6	S19-SLF6	S17-SLF6	PIS2-SLF6	PIS3-SLF6	S5-SLF6	S22-SLF6
S9-SLF6	96.4 %	95.4 %	94.9 %	95.2 %	95.9 %	94.7 %	95.4 %	95.4 %	94.9 %	94.9 %
S0m-SLF6	-	94.1 %	93.6 %	93.9 %	94.4 %	93.1 %	93.9 %	93.9 %	93.1 %	93.6 %
Sm-SLF6		-	93.9 %	94.9 %	95.4 %	93.9 %	94.7 %	94.7 %	94.4 %	93.9 %
S7-SLF6			-	95.9 %	94.4 %	93.4 %	94.1 %	94.6 %	93.9 %	93.6 %
S11-SLF6				-	95.9 %	94.9 %	95.7 %	96.2 %	94.7 %	94.9 %
S19-SLF6					-	95.4 %	96.2 %	96.4 %	95.2 %	94.9 %
S17-SLF6						-	97.7 %	97.7 %	93.9 %	93.6 %
PIS2-SLF6							-	99.0 %	95.2 %	94.4 %
PIS3-SLF6								-	95.2 %	94.4 %
S5-SLF6									-	94.4 %

(g) Type-7 SLFs Max. = 99.2 % Min. = 87.3 % Ave. = 91.9 %

	PIS2-SLF7	S17-SLF7 ψ	S22-SLF7 ψ	Sm-SLF7	S19-SLF7 ψ	S0m-SLF7	S11-SLF7	PIS1-SLF7
S9-SLF7	99.2 %	93.1 %	92.1 %	91.8 %	92.6 %	93.1 %	89.0 %	88.8 %
PIS2-SLF7	-	93.0 %	92.0 %	91.7 %	92.5 %	93.0 %	88.9 %	88.6 %
S17-SLF7 ψ		-	91.8 %	91.3 %	92.1 %	92.3 %	88.5 %	88.1 %
S22-SLF7 ψ			-	91.6 %	91.6 %	91.6 %	87.8 %	87.3 %
Sm-SLF7				-	93.6 %	92.9 %	90.8 %	90.1 %
S19-SLF7 ψ					-	96.4 %	93.4 %	93.0 %
S0m-SLF7						-	92.6 %	92.2 %
S11-SLF7							-	98.4 %

(h) Type-8 SLFs Max. = 100.0 % Min. = 88.2 % Ave. = 93.8 %

	S7-SLF8B	S19-SLF8B	S10-SLF8	S5-SLF8	Sm-SLF8	S9-SLF8	S0m-SLF8	S19-SLF8A	S11-SLF8	S22-SLF8	S17-SLF8A	PIS2-SLF8	S17-SLF8B ψ	Ungrouped S7-FBX1
S7-SLF8A	96.4 %	96.4 %	95.1 %	95.1 %	95.7 %	95.8 %	95.9 %	95.9 %	95.1 %	92.1 %	89.8 %	89.5 %	89.3 %	67.5 %
S7-SLF8B	-	100.0 %	97.7 %	97.7 %	98.2 %	97.7 %	97.5 %	97.2 %	96.4 %	94.4 %	91.3 %	90.9 %	90.5 %	68.6 %
S19-SLF8B		-	97.7 %	97.7 %	98.2 %	97.7 %	97.5 %	97.2 %	96.4 %	94.4 %	91.3 %	90.9 %	90.5 %	68.6 %
S10-SLF8			-	96.7 %	97.5 %	96.6 %	96.4 %	96.4 %	95.7 %	93.1 %	90.6 %	89.6 %	90.0 %	68.1 %
S5-SLF8				-	96.9 %	96.9 %	96.4 %	96.4 %	95.1 %	93.1 %	90.0 %	90.0 %	89.5 %	68.1 %
Sm-SLF8					-	97.1 %	96.4 %	96.4 %	95.7 %	93.6 %	91.1 %	90.4 %	89.8 %	68.8 %
S9-SLF8						-	96.9 %	96.4 %	96.4 %	93.5 %	91.1 %	90.6 %	90.4 %	68.3 %
S0m-SLF8							-	98.5 %	95.7 %	93.6 %	91.3 %	90.4 %	90.5 %	68.9 %
S19-SLF8A								-	95.7 %	93.9 %	91.3 %	90.6 %	90.8 %	68.4 %
S11-SLF8									-	92.1 %	90.6 %	90.1 %	89.3 %	67.8 %
S22-SLF8										-	90.0 %	90.6 %	91.3 %	67.3 %
S17-SLF8A											-	90.3 %	88.2 %	66.0 %
PIS2-SLF8												-	88.7 %	66.0 %
S17-SLF8B ψ													-	67.1 %

Supplementary Table 3 (Continued)

(i) Type-9 SLFs															Max. = 100.0 %		Min. = 92.7 %		Ave. = 95.1 %	
	S17-SLF9A	S7-SLF9A	S9-SLF9A	S11-SLF9	S5-SLF9	Sm-SLF9A	Sm-SLF9B	S22-SLF9	PIS1-SLF9	PIS2-SLF9	S9-SLF9B	S17-SLF9B	S7-SLF9B _y	Ungrouped						
														S0m-FBX1	S19-FBX1					
PIS3-SLF9	97.1 %	97.1 %	97.6 %	95.5 %	94.7 %	95.2 %	94.1 %	94.7 %	94.9 %	94.9 %	94.1 %	95.2 %	95.3 %	87.0 %	87.0 %					
S17-SLF9A	-	97.9 %	98.4 %	96.3 %	94.2 %	95.2 %	94.5 %	95.0 %	95.2 %	95.2 %	94.0 %	95.5 %	96.7 %	87.7 %	86.9 %					
S7-SLF9A		-	99.0 %	96.9 %	94.8 %	95.5 %	95.0 %	95.5 %	95.5 %	95.5 %	94.8 %	96.1 %	97.2 %	87.7 %	87.4 %					
S9-SLF9A			-	97.4 %	95.3 %	96.0 %	95.5 %	96.1 %	96.0 %	96.0 %	95.0 %	96.6 %	97.7 %	88.2 %	88.0 %					
S11-SLF9				-	93.7 %	94.4 %	94.5 %	95.0 %	94.4 %	94.4 %	93.7 %	94.8 %	95.8 %	86.6 %	86.6 %					
S5-SLF9					-	97.3 %	96.1 %	94.0 %	94.1 %	94.1 %	92.7 %	94.0 %	94.9 %	86.6 %	86.1 %					
Sm-SLF9A						-	96.5 %	94.7 %	94.9 %	94.9 %	93.4 %	94.7 %	93.5 %	87.0 %	86.7 %					
Sm-SLF9B							-	94.0 %	93.6 %	93.6 %	93.4 %	94.0 %	94.0 %	86.6 %	86.1 %					
S22-SLF9								-	94.4 %	94.4 %	93.2 %	94.8 %	93.0 %	86.6 %	86.1 %					
PIS1-SLF9									-	100.0 %	93.4 %	94.7 %	94.0 %	86.5 %	85.9 %					
PIS2-SLF9										-	93.4 %	94.7 %	94.0 %	86.5 %	85.9 %					
S9-SLF9B											-	94.5 %	94.0 %	85.6 %	85.6 %					
S17-SLF9B												-	95.8 %	86.9 %	86.9 %					
S7-SLF9B _y													-	91.2 %	90.2 %					
Ungrouped	S0m-FBX1													-	91.9 %					

(j) Type-10 SLFs															Max. = 100.0 %		Min. = 92.0 %		Ave. = 97.1 %	
	PIS3-SLF10	S5-SLF10	S19-SLF10A	S9-SLF10	PIS2-SLF10	S7-SLF10	S11-SLF10	S17-SLF10	S0m-SLF10	S10-SLF10	Sm-SLF10	S22-SLF10	S19-SLF10B	Ungrouped						
														S0m-FBX1	S19-FBX1					
PIS1-SLF10	98.1 %	98.9 %	98.7 %	98.7 %	97.9 %	97.3 %	97.3 %	97.3 %	97.3 %	97.6 %	97.9 %	97.9 %	93.6 %	85.9 %	88.8 %					
PIS3-SLF10	-	97.6 %	97.3 %	97.3 %	96.6 %	96.0 %	96.0 %	96.0 %	96.0 %	96.3 %	96.6 %	96.6 %	92.6 %	84.6 %	87.8 %					
S5-SLF10		-	99.7 %	98.2 %	97.1 %	97.3 %	97.3 %	97.3 %	97.3 %	97.4 %	97.6 %	97.6 %	92.7 %	85.1 %	88.0 %					
S19-SLF10A			-	97.9 %	96.8 %	97.1 %	97.1 %	97.1 %	97.1 %	97.1 %	97.4 %	97.4 %	92.4 %	84.8 %	87.7 %					
S9-SLF10				-	98.2 %	97.3 %	97.3 %	97.3 %	97.3 %	97.6 %	97.9 %	97.9 %	92.9 %	85.8 %	88.5 %					
PIS2-SLF10					-	96.8 %	96.8 %	96.8 %	96.8 %	97.1 %	97.4 %	97.4 %	92.6 %	85.7 %	88.3 %					
S7-SLF10						-	100.0 %	100.0 %	100.0 %	98.9 %	99.2 %	99.2 %	92.0 %	84.9 %	87.5 %					
S11-SLF10							-	100.0 %	100.0 %	98.9 %	99.2 %	99.2 %	92.0 %	84.9 %	87.5 %					
S17-SLF10								-	100.0 %	98.9 %	99.2 %	99.2 %	92.0 %	84.9 %	87.5 %					
S0m-SLF10									-	98.9 %	99.2 %	99.2 %	92.0 %	84.9 %	87.5 %					
S10-SLF10										-	99.7 %	99.7 %	92.0 %	85.2 %	87.5 %					
Sm-SLF10											-	100.0 %	92.3 %	85.4 %	87.8 %					
S22-SLF10												-	92.3 %	85.4 %	87.8 %					
S19-SLF10B													-	86.9 %	88.7 %					
Ungrouped	S0m-FBX1													-	91.9 %					

(k) Type-11 SLFs															Max. = 97.2 %		Min. = 91.2 %		Ave. = 93.7 %	
	Sm-SLF11	S19-SLF11	S10-SLF11	S7-SLF11	S11-SLF11	S22-SLF11	S9-SLF11	S17-SLF11	S0m-SLF11											
S5-SLF11	97.2 %	95.1 %	95.6 %	94.9 %	95.1 %	93.8 %	92.2 %	93.1 %	92.6 %											
Sm-SLF11	-	94.6 %	94.4 %	94.1 %	94.4 %	93.3 %	91.4 %	92.3 %	91.8 %											
S19-SLF11		-	94.4 %	93.6 %	94.4 %	93.3 %	91.9 %	92.6 %	92.8 %											
S10-SLF11			-	93.6 %	94.4 %	93.1 %	91.9 %	92.3 %	93.3 %											
S7-SLF11				-	97.2 %	95.6 %	93.5 %	94.6 %	92.1 %											
S11-SLF11					-	96.7 %	94.5 %	96.2 %	93.1 %											
S22-SLF11						-	93.5 %	94.9 %	91.5 %											
S9-SLF11							-	95.1 %	91.2 %											
S17-SLF11								-	91.3 %											

100%
more than 90 %, less than 100 %
more than 80 %, less than 90 %
more than 70 %, less than 80 %
less than 70 %

100%
more than 90 %, less than 100 %
more than 80 %, less than 90 %
more than 70 %, less than 80 %
less than 70 %

Supplementary Table 3 (Continued)

(l) Type-12 SLFs										Max. = 100.0 %	Min. = 91.9 %	Ave. = 95.9 %
	S0m-SLF12	Sm-SLF12A	S19-SLF12	S10-SLF12	Sm-SLF12B	S9-SLF12ψ	S5-SLF12	S22-SLF12	S17-SLF12B	S17-SLF12Cψ		
S17-SLF12A	100.0 %	100.0 %	97.7 %	96.9 %	96.2 %	95.9 %	97.5 %	94.9 %	96.2 %	92.9 %		
S0m-SLF12	-	100.0 %	97.7 %	96.9 %	96.2 %	95.9 %	97.5 %	94.9 %	96.2 %	92.9 %		
Sm-SLF12A		-	97.7 %	96.9 %	96.2 %	95.9 %	97.5 %	94.9 %	96.2 %	92.9 %		
S19-SLF12			-	97.2 %	96.4 %	96.2 %	96.9 %	95.2 %	96.2 %	93.4 %		
S10-SLF12				-	96.7 %	96.9 %	96.2 %	94.9 %	95.9 %	92.9 %		
Sm-SLF12B					-	96.7 %	95.4 %	94.7 %	95.4 %	92.4 %		
S9-SLF12ψ						-	95.4 %	93.9 %	94.9 %	91.9 %		
S5-SLF12							-	95.2 %	96.7 %	93.4 %		
S22-SLF12								-	97.2 %	95.7 %		
S17-SLF12B									-	94.7 %		

(m) Type-13 SLFs										Max. = 99.7 %	Min. = 91.2 %	Ave. = 95.3 %
	Sm-SLF13	S5-SLF13	S10-SLF13	S17-SLF13	S22-SLF13	S19-SLF13	S0m-SLF13	S9-SLF13	S7-SLF13			
S11-SLF13	96.9 %	94.8 %	95.3 %	95.1 %	95.9 %	95.6 %	96.1 %	94.6 %	92.5 %			
Sm-SLF13	-	95.9 %	94.8 %	94.6 %	96.9 %	96.6 %	95.6 %	94.1 %	91.2 %			
S5-SLF13		-	97.2 %	94.6 %	96.4 %	96.1 %	95.4 %	95.6 %	92.3 %			
S10-SLF13			-	95.4 %	95.6 %	95.4 %	96.6 %	95.4 %	94.1 %			
S17-SLF13				-	97.7 %	97.4 %	96.6 %	95.1 %	92.5 %			
S22-SLF13					-	99.7 %	97.4 %	95.9 %	92.8 %			
S19-SLF13						-	97.2 %	95.6 %	92.5 %			
S0m-SLF13							-	95.9 %	93.6 %			
S9-SLF13								-	92.5 %			

(p) Type-16 SLFs										Max. = 99.0 %	Min. = 90.9 %	Ave. = 94.2 %
	S9-SLF16B	S11-SLF16	S0m-SLF16	S7-SLF16	S22-SLF16	S17-SLF16	S5-SLF16	S10-SLF16	Sm-SLF16	S19-SLF16		
S9-SLF16A	99.0 %	97.4 %	95.9 %	93.9 %	92.3 %	92.3 %	92.4 %	97.4 %	98.4 %	92.7 %		
S9-SLF16B	-	96.9 %	95.7 %	93.4 %	91.8 %	91.8 %	91.9 %	96.9 %	94.0 %	92.2 %		
S11-SLF16		-	96.4 %	94.4 %	92.8 %	92.8 %	93.0 %	98.0 %	93.2 %	93.0 %		
S0m-SLF16			-	94.1 %	92.8 %	92.8 %	93.0 %	96.9 %	95.6 %	93.5 %		
S7-SLF16				-	92.6 %	91.8 %	90.9 %	95.1 %	94.8 %	91.5 %		
S22-SLF16					-	96.4 %	93.2 %	93.9 %	95.0 %	94.5 %		
S17-SLF16						-	92.4 %	94.4 %	94.5 %	94.3 %		
S5-SLF16							-	94.0 %	95.3 %	94.8 %		
S10-SLF16								-	91.5 %	94.0 %		
Sm-SLF16									-	98.4 %		

(n) Type-14 SLFs										Max. = 97.4 %	Min. = 93.6 %	Ave. = 95.2 %
	S7-SLF14	S19-SLF14	S0m-SLF14	S9-SLF14	S22-SLF14	S11-SLF14	S17-SLF14	Sm-SLF14				
S10-SLF14	97.4 %	95.7 %	93.8 %	94.1 %	93.8 %	95.4 %	94.6 %	93.6 %				
S7-SLF14	-	96.3 %	94.8 %	95.6 %	94.8 %	95.4 %	95.1 %	94.6 %				
S19-SLF14		-	95.6 %	95.6 %	95.3 %	95.6 %	94.8 %	95.0 %				
S0m-SLF14			-	96.9 %	95.6 %	96.7 %	95.1 %	94.6 %				
S9-SLF14				-	95.6 %	97.2 %	95.4 %	94.3 %				
S22-SLF14					-	95.9 %	94.1 %	94.1 %				
S11-SLF14						-	96.9 %	94.1 %				
S17-SLF14							-	94.1 %				

(o) Type-15 SLFs							Max. = 99.0 %	Min. = 93.4 %	Ave. = 96.3 %
	S5-SLF15ψ	S11-SLF15ψ	S0m-SLF15ψ	S22-SLF15	S19-SLF15ψ	S9-SLF15			
Sm-SLF15	98.5 %	98.2 %	98.2 %	98.0 %	95.2 %	93.6 %			
S5-SLF15ψ	-	98.2 %	98.2 %	98.0 %	95.2 %	93.4 %			
S11-SLF15ψ		-	99.0 %	97.7 %	95.5 %	93.9 %			
S0m-SLF15ψ			-	98.2 %	95.5 %	93.6 %			
S22-SLF15				-	95.7 %	93.9 %			
S19-SLF15ψ					-	95.2 %			

(q) Type-17 SLFs						Max. = 100.0 %	Min. = 99.7 %	Ave. = 99.9 %
	S9-SLF17	S10-SLF17	S17-SLF17	S0m-SLF17	S5-SLF17			
S7-SLF17	100.0 %	100.0 %	100.0 %	100.0 %	99.7 %			
S9-SLF17	-	100.0 %	100.0 %	100.0 %	99.7 %			
S10-SLF17		-	100.0 %	100.0 %	99.7 %			
S17-SLF17			-	100.0 %	99.7 %			
S0m-SLF17				-	99.7 %			

(r) Type-18 SLFs				
	S10-SLF18	S22-SLF18	Sm-SLF18	
S5-SLF18	100.0 %	100.0 %	100.0 %	100%
S10-SLF18	-	100.0 %	100.0 %	more than 90 %, less than 100 %
S22-SLF18		-	100.0 %	more than 80 %, less than 90 %
				more than 70 %, less than 80 %
				less than 70 %

Supplementary Table 3 (Continued)

(s) SLFs of S5-haplotype		Max. = 92.5 %		Min. = 44.2 %		Ave. = 52.0 %											
	S5-SLF2	S5-SLF3	S5-SLF4	S5-SLF5	S5-SLF5B	S5-SLF6	S5-SLF8	S5-SLF9	S5-SLF10	S5-SLF11	S5-SLF12	S5-SLF13	S5-SLF15ψ	S5-SLF16	S5-SLF17	S5-SLF18	S5-FBX2
S5-SLF1	58.4 %	51.0 %	47.2 %	53.4 %	52.9 %	48.5 %	55.0 %	52.1 %	49.5 %	54.1 %	48.5 %	49.5 %	52.2 %	52.0 %	52.6 %	47.6 %	50.8 %
S5-SLF2	-	52.1 %	51.8 %	53.6 %	52.3 %	49.5 %	54.3 %	53.3 %	51.3 %	53.0 %	51.3 %	50.4 %	50.0 %	51.3 %	50.9 %	46.8 %	54.6 %
S5-SLF3	-	-	51.7 %	50.9 %	49.3 %	47.3 %	52.2 %	51.9 %	51.2 %	54.9 %	50.3 %	70.3 %	55.6 %	54.1 %	54.5 %	47.8 %	56.0 %
S5-SLF4			-	56.8 %	55.1 %	53.4 %	48.7 %	48.3 %	46.4 %	45.7 %	75.6 %	47.0 %	48.9 %	48.6 %	50.0 %	50.0 %	51.3 %
S5-SLF5				-	92.5 %	53.0 %	50.3 %	50.6 %	48.6 %	48.5 %	56.7 %	48.0 %	51.7 %	50.1 %	50.3 %	52.9 %	51.5 %
S5-SLF5B					-	51.8 %	48.3 %	51.2 %	48.8 %	47.8 %	56.5 %	46.4 %	50.4 %	49.9 %	51.3 %	50.9 %	50.1 %
S5-SLF6						-	47.3 %	47.5 %	47.4 %	47.6 %	52.7 %	44.4 %	49.2 %	47.7 %	47.8 %	59.0 %	47.8 %
S5-SLF8							-	51.0 %	48.0 %	53.6 %	49.2 %	51.4 %	53.6 %	55.7 %	53.9 %	47.4 %	53.4 %
S5-SLF9								-	89.0 %	49.6 %	45.5 %	49.5 %	53.8 %	51.3 %	50.9 %	44.8 %	54.3 %
S5-SLF10									-	48.1 %	44.2 %	48.2 %	54.6 %	50.3 %	49.9 %	44.2 %	53.8 %
S5-SLF11										-	49.4 %	53.4 %	51.7 %	50.0 %	49.6 %	45.3 %	54.1 %
S5-SLF12											-	46.4 %	48.6 %	47.8 %	48.2 %	48.5 %	49.2 %
S5-SLF13												-	51.5 %	52.2 %	52.9 %	45.9 %	52.2 %
S5-SLF15ψ													-	63.9 %	62.4 %	46.7 %	63.1 %
S5-SLF16														-	79.2 %	45.5 %	70.7 %
S5-SLF17															-	44.5 %	67.9 %
S5-SLF18																-	47.1 %

(t) SLFs of S7-haplotype		Max. = 97.2 %		Min. = 45.4 %		Ave. = 53.4 %											
	S7-SLF1B	S7-SLF2	S7-SLF3	S7-SLF4	S7-SLF5	S7-SLF6	S7-SLF8A	S7-SLF8B	S7-SLF9A	S7-SLF9Bψ	S7-SLF10	S7-SLF11	S7-SLF13	S7-SLF14	S7-SLF16	S7-SLF17	S7-FBX1
S7-SLF1	95.1 %	58.4 %	49.9 %	47.7 %	53.2 %	49.5 %	54.1 %	54.6 %	51.9 %	51.2 %	49.2 %	53.6 %	49.2 %	53.1 %	52.1 %	51.6 %	53.8 %
S7-SLF1B	-	59.6 %	51.0 %	48.5 %	54.6 %	50.3 %	53.6 %	54.1 %	52.1 %	50.7 %	49.1 %	53.2 %	49.9 %	52.9 %	51.8 %	51.0 %	53.3 %
S7-SLF2		-	50.9 %	51.5 %	51.4 %	48.4 %	52.4 %	53.2 %	52.9 %	51.4 %	50.5 %	51.8 %	50.3 %	53.3 %	53.5 %	52.0 %	52.3 %
S7-SLF3			-	51.8 %	53.7 %	49.0 %	53.5 %	53.8 %	51.8 %	50.9 %	50.4 %	53.9 %	66.0 %	52.5 %	49.9 %	52.0 %	52.5 %
S7-SLF4				-	58.7 %	53.7 %	49.1 %	50.1 %	49.2 %	48.8 %	48.2 %	47.0 %	48.6 %	50.0 %	51.2 %	50.8 %	50.4 %
S7-SLF5					-	52.3 %	49.6 %	50.0 %	50.6 %	52.5 %	48.1 %	47.7 %	49.1 %	50.5 %	50.3 %	50.0 %	53.9 %
S7-SLF6						-	46.2 %	47.8 %	48.2 %	48.2 %	47.2 %	47.3 %	45.4 %	47.6 %	46.7 %	46.2 %	47.8 %
S7-SLF8A							-	96.4 %	50.3 %	47.3 %	47.2 %	52.4 %	52.5 %	53.0 %	54.6 %	53.1 %	67.5 %
S7-SLF8B								-	51.3 %	48.6 %	47.7 %	53.6 %	52.9 %	54.0 %	56.1 %	54.7 %	68.6 %
S7-SLF9A									-	97.2 %	88.3 %	50.9 %	51.8 %	54.7 %	51.6 %	51.2 %	52.2 %
S7-SLF9Bψ										-	88.4 %	49.3 %	47.7 %	53.4 %	48.4 %	47.8 %	49.8 %
S7-SLF10											-	48.1 %	48.8 %	52.9 %	49.7 %	49.5 %	49.7 %
S7-SLF11												-	53.4 %	51.5 %	48.5 %	48.3 %	54.6 %
S7-SLF13													-	53.5 %	52.3 %	53.7 %	54.3 %
S7-SLF14														-	75.8 %	72.8 %	53.9 %
S7-SLF16															-	78.5 %	55.6 %
S7-SLF17																-	54.7 %

100%
more than 90 %, less than 100 %
more than 80 %, less than 90 %
more than 70 %, less than 80 %
less than 70 %

Supplementary Table 3 (Continued)

(u) SLFs of S9-haplotype		Max. = 99.0 %		Min. = 43.9 %		Ave. = 53.2 %														
	S9-SLF3	S9-SLF4	S9-SLF5	S9-SLF6	S9-SLF7	S9-SLF8	S9-SLF9A	S9-SLF9B	S9-SLF10	S9-SLF11	S9-SLF12ψ	S9-SLF13	S9-SLF14	S9-SLF15	S9-SLF16A	S9-SLF16B	S9-SLF17	S9-FBX1		
S9-SLF1	50.3 %	46.0 %	52.9 %	49.7 %	55.2 %	55.1 %	51.7 %	50.6 %	49.4 %	51.7 %	46.6 %	47.2 %	53.1 %	54.6 %	51.0 %	50.5 %	51.0 %	52.6 %		
S9-SLF3	-	50.6 %	50.0 %	46.5 %	53.7 %	52.6 %	53.9 %	53.2 %	51.7 %	53.8 %	49.2 %	69.0 %	55.7 %	55.0 %	54.0 %	52.9 %	53.5 %	51.2 %		
S9-SLF4		-	57.1 %	53.7 %	49.6 %	49.6 %	48.4 %	48.1 %	46.5 %	45.5 %	78.2 %	47.8 %	48.4 %	49.1 %	49.2 %	48.7 %	50.0 %	50.9 %		
S9-SLF5			-	52.8 %	51.7 %	50.1 %	50.9 %	49.7 %	48.2 %	48.8 %	56.7 %	47.7 %	50.8 %	51.3 %	49.7 %	49.2 %	50.7 %	52.7 %		
S9-SLF6				-	49.4 %	47.2 %	47.7 %	46.6 %	47.4 %	46.6 %	51.4 %	43.9 %	48.2 %	50.1 %	46.5 %	46.3 %	47.5 %	53.4 %		
S9-SLF7					-	50.6 %	50.8 %	50.9 %	50.9 %	50.9 %	49.9 %	49.5 %	63.6 %	60.2 %	60.5 %	60.0 %	58.9 %	52.3 %		
S9-SLF8						-	51.6 %	50.8 %	49.5 %	52.4 %	50.4 %	51.0 %	54.1 %	54.2 %	56.4 %	56.1 %	54.2 %	51.4 %		
S9-SLF9A							-	95.0 %	90.0 %	50.8 %	45.7 %	51.8 %	54.4 %	55.1 %	51.4 %	50.9 %	51.7 %	49.6 %		
S9-SLF9B								-	87.1 %	49.2 %	45.2 %	51.0 %	53.9 %	53.5 %	50.4 %	49.9 %	50.6 %	49.6 %		
S9-SLF10									-	49.2 %	44.2 %	49.1 %	52.6 %	54.3 %	49.6 %	49.3 %	50.1 %	49.2 %		
S9-SLF11										-	45.6 %	51.7 %	52.6 %	51.4 %	50.0 %	49.2 %	49.5 %	51.9 %		
S9-SLF12ψ											-	48.2 %	47.7 %	48.2 %	47.7 %	47.2 %	48.8 %	50.8 %		
S9-SLF13												-	51.7 %	51.4 %	51.5 %	50.5 %	52.6 %	48.8 %		
S9-SLF14													-	65.1 %	75.5 %	75.0 %	73.3 %	51.5 %		
S9-SLF15														-	65.0 %	64.5 %	63.1 %	53.3 %		
S9-SLF16A															-	99.0 %	78.9 %	49.9 %		
S9-SLF16B																-	78.4 %	49.4 %		
S9-SLF17																	-	50.0 %		

(v) SLFs of S11-haplotype		Max. = 87.2 %		Min. = 44.4 %		Ave. = 52.3 %														
	S11-SLF2	S11-SLF3	S11-SLF3B	S11-SLF4	S11-SLF5	S11-SLF6	S11-SLF7	S11-SLF8	S11-SLF9	S11-SLF10	S11-SLF11	S11-SLF13	S11-SLF14	S11-SLF15ψ	S11-SLF16	S11-FBX1				
S11-SLF1	56.8 %	49.6 %	50.9 %	47.0 %	52.8 %	50.5 %	54.5 %	53.6 %	50.8 %	47.8 %	52.7 %	49.9 %	51.9 %	51.5 %	49.5 %	50.1 %	100 %			
S11-SLF2	-	52.8 %	51.8 %	48.5 %	51.4 %	47.8 %	54.4 %	52.7 %	52.6 %	49.0 %	54.4 %	50.1 %	53.9 %	49.1 %	53.7 %	47.5 %	more than 100 %			
S11-SLF3		-	74.4 %	50.1 %	49.9 %	47.0 %	53.5 %	53.2 %	52.6 %	50.8 %	53.8 %	69.5 %	57.6 %	54.3 %	54.5 %	47.7 %	more than 100 %			
S11-SLF3B			-	50.5 %	50.0 %	48.6 %	54.9 %	55.2 %	53.0 %	51.7 %	54.7 %	67.4 %	53.2 %	53.9 %	50.1 %	48.3 %	more than 100 %			
S11-SLF4				-	56.6 %	53.2 %	50.0 %	50.0 %	47.8 %	46.3 %	46.7 %	47.3 %	48.2 %	48.9 %	49.0 %	54.3 %	less than 70 %			
S11-SLF5					-	52.8 %	50.4 %	51.0 %	50.9 %	48.3 %	48.0 %	48.8 %	50.6 %	51.4 %	49.5 %	54.0 %				
S11-SLF6						-	48.6 %	48.5 %	48.4 %	47.2 %	48.9 %	44.4 %	49.7 %	50.0 %	46.0 %	64.3 %				
S11-SLF7							-	50.4 %	52.3 %	51.3 %	51.3 %	50.9 %	63.5 %	58.7 %	58.5 %	48.8 %				
S11-SLF8								-	52.1 %	49.3 %	54.1 %	53.4 %	54.9 %	53.1 %	55.9 %	49.6 %				
S11-SLF9									-	87.2 %	49.9 %	50.9 %	56.3 %	54.7 %	51.7 %	47.8 %				
S11-SLF10										-	48.7 %	48.8 %	54.1 %	54.8 %	49.7 %	46.0 %				
S11-SLF11											-	53.6 %	53.1 %	51.4 %	49.6 %	47.4 %				
S11-SLF13												-	52.9 %	50.9 %	51.0 %	47.4 %				
S11-SLF14													-	65.7 %	76.0 %	47.5 %				
S11-SLF15ψ														-	64.4 %	51.7 %				
S11-SLF16															-	45.9 %				

100%
more than 90 %, less than 100 %
more than 80 %, less than 90 %
more than 70 %, less than 80 %
less than 70 %

Supplementary Table 3 (Continued)

(w) SLFs of S17-haplotype	Max. = 96.2 %			Min. = 43.2 %			Ave. = 52.9 %															
	S17-SLF1B	S17-SLF2	S17-SLF3	S17-SLF4	S17-SLF4Bψ	S17-SLF5	S17-SLF6	S17-SLF7ψ	S17-SLF8A	S17-SLF8Bψ	S17-SLF9A	S17-SLF9B	S17-SLF10	S17-SLF11	S17-SLF12A	S17-SLF12B	S17-SLF12Cψ	S17-SLF13	S17-SLF14	S17-SLF16	S17-SLF17	
S17-SLF1	87.9 %	58.6 %	50.1 %	48.1 %	47.0 %	53.3 %	49.7 %	53.9 %	51.9 %	50.4 %	48.7 %	50.0 %	46.8 %	50.4 %	46.6 %	47.1 %	45.3 %	49.1 %	51.4 %	51.0 %	49.5 %	
S17-SLF1B	-	56.6 %	51.5 %	49.0 %	47.7 %	53.7 %	52.0 %	56.6 %	54.0 %	52.3 %	49.9 %	51.2 %	47.9 %	50.8 %	47.6 %	48.4 %	46.3 %	49.6 %	52.1 %	52.0 %	50.8 %	
S17-SLF2	-	-	49.9 %	49.0 %	48.6 %	52.0 %	48.6 %	54.4 %	51.7 %	50.6 %	53.2 %	53.7 %	51.7 %	51.0 %	49.6 %	49.3 %	48.2 %	50.1 %	53.0 %	51.6 %	51.2 %	
S17-SLF3			-	50.7 %	50.9 %	51.2 %	48.2 %	53.5 %	52.5 %	51.9 %	52.5 %	53.5 %	51.6 %	53.4 %	50.3 %	50.8 %	50.0 %	70.6 %	56.0 %	54.0 %	54.5 %	
S17-SLF4				-	94.0 %	56.6 %	53.4 %	48.2 %	49.1 %	48.6 %	47.6 %	47.6 %	46.7 %	45.4 %	74.8 %	75.1 %	73.0 %	46.9 %	47.7 %	49.0 %	49.2 %	
S17-SLF4Bψ					-	57.4 %	52.9 %	49.9 %	48.9 %	48.3 %	48.2 %	48.5 %	47.3 %	45.5 %	75.6 %	75.6 %	73.6 %	48.2 %	47.7 %	50.0 %	50.0 %	
S17-SLF5						-	53.0 %	50.4 %	50.5 %	49.9 %	51.4 %	51.3 %	49.6 %	48.0 %	57.0 %	57.2 %	54.2 %	49.3 %	50.1 %	51.0 %	50.4 %	
S17-SLF6							-	49.0 %	47.6 %	48.1 %	47.7 %	48.5 %	47.8 %	46.3 %	50.6 %	51.1 %	49.9 %	44.1 %	49.0 %	48.3 %	47.0 %	
S17-SLF7ψ								-	51.0 %	50.5 %	51.8 %	52.0 %	50.4 %	50.6 %	49.0 %	49.2 %	49.2 %	51.0 %	63.3 %	61.8 %	58.9 %	
S17-SLF8A									-	88.2 %	51.0 %	50.5 %	47.8 %	51.0 %	49.3 %	50.1 %	47.5 %	51.7 %	52.1 %	54.8 %	53.9 %	
S17-SLF8Bψ										-	51.3 %	50.1 %	48.7 %	52.0 %	48.0 %	49.0 %	47.2 %	50.5 %	52.3 %	55.2 %	53.8 %	
S17-SLF9A											-	95.5 %	88.0 %	49.4 %	44.7 %	45.2 %	44.7 %	50.0 %	53.9 %	51.9 %	50.9 %	
S17-SLF9B												-	87.3 %	49.6 %	45.2 %	45.7 %	44.7 %	51.5 %	54.2 %	52.7 %	50.6 %	
S17-SLF10													-	47.4 %	43.5 %	43.5 %	43.2 %	48.6 %	52.8 %	51.6 %	49.5 %	
S17-SLF11														-	46.1 %	45.8 %	45.6 %	51.3 %	51.2 %	48.8 %	49.1 %	
S17-SLF12A															-	96.2 %	92.9 %	47.8 %	48.1 %	48.0 %	48.0 %	
S17-SLF12B																-	94.7 %	47.5 %	48.7 %	48.7 %	48.5 %	
S17-SLF12Cψ																	-	47.8 %	47.7 %	47.4 %	48.6 %	
S17-SLF13																		-	51.1 %	50.8 %	52.6 %	
S17-SLF14																			-	76.8 %	73.0 %	
S17-SLF16																				-	78.6 %	
(x) SLFs of S19-haplotype	Max. = 100.0 %			Min. = 43.5 %			Ave. = 53.7 %															
	S19-SLF1B	S19-SLF2	S19-SLF3	S19-SLF4	S19-SLF5	S19-SLF5B	S19-SLF6	S19-SLF7ψ	S19-SLF8A	S19-SLF8B	S19-SLF10A	S19-SLF10B	S19-SLF11	S19-SLF12	S19-SLF13	S19-SLF14	S19-SLF15ψ	S19-SLF16	S19-FBX1	S19-FBX2	S19-FBX3	
S19-SLF1	97.7 %	60.4 %	50.3 %	48.2 %	53.7 %	53.2 %	50.3 %	55.4 %	55.2 %	54.6 %	49.5 %	51.6 %	53.3 %	48.1 %	49.5 %	54.2 %	54.5 %	52.2 %	51.7 %	51.4 %	51.4 %	
S19-SLF1B	-	60.9 %	51.5 %	48.7 %	53.9 %	53.4 %	50.3 %	56.2 %	55.8 %	55.1 %	49.2 %	51.6 %	54.1 %	48.1 %	50.0 %	55.2 %	54.7 %	53.3 %	51.7 %	51.7 %	51.7 %	
S19-SLF2		-	52.7 %	50.9 %	51.7 %	50.9 %	48.8 %	54.9 %	53.8 %	53.9 %	51.4 %	53.0 %	53.4 %	50.9 %	51.7 %	54.6 %	51.8 %	54.1 %	52.9 %	53.4 %	52.4 %	
S19-SLF3			-	51.4 %	50.1 %	49.6 %	48.1 %	54.0 %	53.4 %	51.9 %	50.6 %	52.5 %	55.7 %	50.3 %	69.0 %	56.4 %	54.7 %	54.4 %	50.9 %	51.2 %	52.6 %	
S19-SLF4				-	56.8 %	55.4 %	54.2 %	49.9 %	49.1 %	49.2 %	45.4 %	46.8 %	46.0 %	74.0 %	46.3 %	47.7 %	48.2 %	50.1 %	47.4 %	48.0 %	58.2 %	
S19-SLF5					-	92.5 %	54.1 %	50.1 %	50.5 %	49.7 %	48.1 %	48.1 %	48.0 %	56.7 %	48.8 %	51.4 %	52.4 %	50.7 %	49.2 %	51.7 %	57.7 %	
S19-SLF5B						-	52.8 %	50.4 %	49.1 %	48.3 %	48.8 %	49.6 %	48.6 %	57.0 %	47.0 %	52.0 %	51.4 %	51.4 %	49.5 %	52.4 %	57.7 %	
S19-SLF6							-	49.1 %	48.3 %	47.1 %	47.4 %	47.9 %	48.1 %	50.9 %	44.6 %	48.8 %	50.6 %	47.8 %	48.7 %	50.3 %	53.7 %	
S19-SLF7ψ								-	51.3 %	51.1 %	51.5 %	52.7 %	53.3 %	49.6 %	51.4 %	63.9 %	59.4 %	61.7 %	52.7 %	54.2 %	54.5 %	
S19-SLF8A									-	97.2 %	49.4 %	49.6 %	53.2 %	49.4 %	52.6 %	55.4 %	53.8 %	56.5 %	49.6 %	51.4 %	52.5 %	
S19-SLF8B										-	48.3 %	49.6 %	52.9 %	49.9 %	52.2 %	54.9 %	53.8 %	57.0 %	49.2 %	51.8 %	52.5 %	
S19-SLF10A											-	92.1 %	48.8 %	43.5 %	48.7 %	53.5 %	54.7 %	50.8 %	87.7 %	76.6 %	50.3 %	
S19-SLF10B												-	51.7 %	45.9 %	50.5 %	54.5 %	56.1 %	53.4 %	88.7 %	77.1 %	51.3 %	
S19-SLF11													-	48.9 %	53.8 %	53.5 %	50.8 %	51.4 %	49.4 %	51.9 %	50.3 %	
S19-SLF12														-	47.5 %	49.4 %	47.2 %	49.1 %	43.8 %	46.0 %	57.0 %	
S19-SLF13															-	52.5 %	50.9 %	52.5 %	48.7 %	49.7 %	49.0 %	
S19-SLF14																-	66.1 %	76.9 %	54.5 %	53.8 %	54.2 %	
S19-SLF15ψ																	-	65.9 %	54.4 %	55.6 %	52.8 %	
S19-SLF16																		-	52.1 %	52.1 %	52.5 %	
S19-FBX1																			-	80.2 %	52.2 %	
S19-FBX2																				-	52.8 %	

Supplementary Table 3 (Continued)

(y) S-RNases	Max. = 94.1 %		Min. = 35.2 %		Ave. = 52.9 %										
	PIS2-RNase	PIS13-RNase	PIS3-RNase	PIS12-RNase	PIS1-RNase	S10-RNase	S17-RNase	S9-RNase	PIS7-RNase	S7-RNase	S11-RNase	S5-RNase	Sm-RNase	S19-RNase	S0m-RNase
S22-RNase	94.1 %	85.3 %	81.7 %	76.8 %	71.1 %	60.6 %	61.0 %	53.2 %	54.7 %	43.4 %	42.9 %	43.6 %	39.9 %	43.4 %	42.6 %
PIS2-RNase	-	86.5 %	80.2 %	77.4 %	71.2 %	63.1 %	60.4 %	53.6 %	54.7 %	45.2 %	42.0 %	45.5 %	40.9 %	43.3 %	41.1 %
PIS13-RNase		-	81.6 %	76.7 %	70.6 %	60.1 %	54.9 %	50.6 %	54.7 %	36.4 %	35.2 %	42.4 %	39.1 %	40.5 %	39.7 %
PIS3-RNase			-	84.0 %	72.6 %	63.2 %	62.5 %	55.4 %	54.7 %	45.7 %	43.0 %	45.1 %	40.9 %	44.4 %	42.7 %
PIS12-RNase				-	75.0 %	65.0 %	58.5 %	52.8 %	57.5 %	39.6 %	35.8 %	41.3 %	38.8 %	40.0 %	39.5 %
PIS1-RNase					-	65.0 %	58.3 %	55.6 %	56.3 %	44.8 %	41.2 %	44.7 %	39.9 %	44.0 %	41.5 %
S10-RNase						-	68.0 %	60.2 %	59.0 %	45.4 %	43.1 %	49.8 %	41.2 %	43.6 %	43.1 %
S17-RNase							-	57.3 %	54.0 %	47.0 %	45.7 %	47.1 %	42.7 %	45.9 %	47.5 %
S9-RNase								-	84.9 %	47.2 %	45.5 %	47.2 %	43.8 %	45.0 %	44.5 %
PIS7-RNase									-	43.3 %	39.1 %	41.3 %	40.5 %	41.5 %	39.6 %
S7-RNase										-	68.7 %	51.6 %	46.9 %	47.5 %	47.5 %
S11-RNase											-	50.2 %	44.5 %	48.9 %	48.9 %
S5-RNase												-	47.1 %	46.2 %	47.1 %
Sm-RNase													-	72.8 %	73.8 %
S19-RNase														-	86.5 %

100%
 more than 90 %, less than 100 %
 more than 80 %, less than 90 %
 more than 70 %, less than 80 %
 less than 70 %

Pairwise comparisons among allelic *SLFs* of the same type (a–r), among *SLFs* of each S-haplotype (s–x), or among allelic *S-RNases* (y). Shading shows the level of identity. Each value was computed by using the GENETYX-MAC (ver. 16.0.6).

Supplementary Table 4 Summary of transformation experiments

Plant ID	S-genotype ^a		Transgene expression ^b	self pollination	
				seed set ^c	Phenotype
S7-SLF1 #41	<i>S5S22</i>	<i>S11S22</i> x #14 (<i>S5S17/S7-SLF1</i>) ^d	+++	+++	SC
S7-SLF1 #42	<i>S11S19</i>	<i>S7S11</i> x #37 (<i>S5S19/S7-SLF1</i>) ^d	+++	-	SI
S7-SLF1 #43	<i>S11S19</i>	<i>S7S11</i> x #40 (<i>S17S19/S7-SLF1</i>) ^d	+++	-	SI
S7-SLF1 #44	<i>S11S22</i>	#42 x <i>S11S22</i>	+++	+++	SC
S7-SLF1 #45	<i>S19S22</i>	#42 x <i>S11S22</i>	+++	+++	SC
S7-SLF1 #46	<i>S19S22</i>	#43 x <i>S11S22</i>	+++	+++	SC
S5-SLF3 #1	<i>S5S11</i>	T0	+++	-	SI
S5-SLF3 #2	<i>S5S17</i>	T0	+++	-	SI
S5-SLF3 #3	<i>S11S7</i>	<i>S7S7</i> x #1	++	+++	SC
S5-SLF3 #4	<i>S11S7</i>	<i>S7S7</i> x #1	++	+++	SC
S5-SLF3 #5	<i>S9S11</i>	<i>S9S9</i> x #1	+	-	SI
S5-SLF3 #6	<i>S5S17</i>	<i>S17S17</i> x #1	+++	-	SI
S5-SLF3 #7	<i>S5S17</i>	<i>S17S17</i> x #1	+++	-	SI
S5-SLF3 #8	<i>S5S19</i>	<i>S19S19</i> x #1	+	-	SI
S5-SLF3 #9	<i>S5S9</i>	<i>S9S11</i> x #2	+++	-	SI
S5-SLF3 #10	<i>S5S11</i>	<i>S9S11</i> x #2	+++	-	SI
S5-SLF3 #11	<i>S5S11</i>	<i>S9S11</i> x #2	+++	-	SI
S5-SLF3 #12	<i>S9S17</i>	<i>S9S11</i> x #2	+++	-	SI
S5-SLF3 #13	<i>S5S7</i>	<i>S7S7</i> x #2	+++	+++	SC
S5-SLF3 #14	<i>S5S7</i>	<i>S7S7</i> x #2	+++	+++	SC
S5-SLF3 #15	<i>S5S19</i>	<i>S19S19</i> x #2	+++	-	SI
S5-SLF3 #16	<i>S5S19</i>	<i>S19S19</i> x #2	+++	-	SI
S7-SLF3 #1	<i>S5S7</i>	T0	+++	-	SI
S7-SLF3 #2	<i>S5S7</i>	T0	+++	-	SI
S7-SLF3 #3	<i>S5S7</i>	T0	+++	-	SI
S7-SLF3 #4	<i>S5S7</i>	T0	+++	-	SI
S7-SLF3 #5	<i>S5S7</i>	T0	+++	-	SI
S7-SLF3 #6	<i>S5S17</i>	T0	+++	-	SI
S7-SLF3 #7	<i>S5S17</i>	T0	+++	-	SI
S7-SLF3 #8	<i>S5S17</i>	T0	+++	-	SI
S7-SLF3 #9	<i>S7S9</i>	T0	+++	-	SI
S7-SLF3 #10	<i>S7S9</i>	T0	++	-	SI
S7-SLF3 #11	<i>S7S9</i>	T0	++	-	SI
S7-SLF3 #12	<i>S7S9</i>	T0	+++	-	SI
S7-SLF3 #13	<i>S7S11</i>	#1 x <i>S11S11</i>	+++	-	SI
S7-SLF3 #14	<i>S7S11</i>	#2 x <i>S11S11</i>	+++	-	SI
S7-SLF3 #15	<i>S7S11</i>	#3 x <i>S11S11</i>	+++	-	SI
S7-SLF3 #16	<i>S7S19</i>	#1 x <i>S5S19</i>	+++	-	SI
S7-SLF3 #17	<i>S7S19</i>	#3 x <i>S5S19</i>	+++	-	SI
S7-SLF3 #18	<i>S5S19</i>	#6 x <i>S5S19</i>	+++	-	SI
S11-SLF3B #1	<i>S5S17</i>	T0	+++	-	SI
S11-SLF3B #2	<i>S5S17</i>	T0	+++	-	SI
S11-SLF3B #3	<i>S5S17</i>	T0	+++	-	SI
S11-SLF3B #4	<i>S5S11</i>	T0	++	-	SI
S11-SLF3B #5	<i>S5S11</i>	T0	+++	-	SI

S11-SLF3B #6	S5S9	S9S11 x #1	+++	-	SI
S11-SLF3B #7	S5S9	S9S11 x #1	+++	-	SI
S11-SLF3B #8	S5S11	S9S11 x #1	+++	-	SI
S11-SLF3B #9	S5S11	S9S11 x #1	+++	-	SI
S11-SLF3B #10	S11S17	S11S11 x #1	+++	-	SI
S11-SLF3B #11	S11S17	S11S11 x #1	+++	-	SI
S11-SLF3B #12	S5S7	S7S19 x #1	+++	-	SI
S11-SLF3B #13	S5S19	S7S19 x #1	+++	-	SI
S11-SLF3B #14	S5S19	S7S19 x #1	+++	-	SI
S11-SLF3B #15	S5S7	S7S19 x #1	++	-	SI
S11-SLF3B #16	S5S19	S7S19 x #1	+++	-	SI
S11-SLF3B #17	S7S17	S7S19 x #1	+++	-	SI
S11-SLF3B #18	S5S9	S9S11 x #3	++	-	SI
S11-SLF3B #19	S9S17	S9S11 x #3	++	-	SI
S11-SLF3B #20	S9S17	S9S11 x #3	++	-	SI
S11-SLF3B #21	S11S17	S9S11 x #3	+++	-	SI
S11-SLF3B #22	S5S7	S7S19 x #3	+++	-	SI
S11-SLF3B #23	S7S17	S7S19 x #3	+++	-	SI
S11-SLF3B #24	S17S19	S7S19 x #3	++	-	SI

S7-SLF9 #1	S5S17	T0	+++	-	SI
S7-SLF9 #2	S5S17	T0	+++	-	SI
S7-SLF9 #3	S5S17	T0	+++	-	SI
S7-SLF9 #4	S5S9	S9S11 x #1	+++	-	SI
S7-SLF9 #5	S5S11	S9S11 x #1	+++	-	SI
S7-SLF9 #6	S5S11	S9S11 x #1	++	-	SI
S7-SLF9 #7	S5S7	S7S19 x #1	+++	-	SI
S7-SLF9 #8	S5S7	S7S19 x #1	+++	-	SI
S7-SLF9 #9	S7S17	S7S19 x #1	++	-	SI
S7-SLF9 #10	S5S19	S7S19 x #1	+++	+++	SC
S7-SLF9 #11	S17S19	S7S19 x #1	+++	+++	SC
S7-SLF9 #12	S5S9	S9S11 x #2	+++	-	SI
S7-SLF9 #13	S5S11	S9S11 x #2	+++	-	SI
S7-SLF9 #14	S11S17	S9S11 x #2	+++	-	SI
S7-SLF9 #15	S5S7	S7S19 x #2	+++	-	SI
S7-SLF9 #16	S5S7	S7S19 x #2	+++	-	SI
S7-SLF9 #17	S5S19	S7S19 x #2	+++	+++	SC
S7-SLF9 #18	S17S19	S7S19 x #2	+++	+++	SC
S7-SLF9 #19	S5S9	S9S11 x #3	+++	-	SI
S7-SLF9 #20	S5S9	S9S11 x #3	+++	-	SI
S7-SLF9 #21	S11S17	S9S11 x #3	+++	-	SI
S7-SLF9 #22	S5S7	S7S19 x #3	+++	-	SI
S7-SLF9 #23	S5S7	S7S19 x #3	+++	-	SI
S7-SLF9 #24	S5S19	S7S19 x #3	+++	+++	SC
S7-SLF9 #25	S5S19	S7S19 x #3	+++	+++	SC
S7-SLF9 #26	S7S9	S7S7 x #4	++	-	SI
S7-SLF9 #27	S7S11	S7S7 x #5	++	-	SI
S7-SLF9 #28	S7S17	S7S7 x #11	+++	-	SI
S7-SLF9 #29	S7S19	S7S7 x #11	++	+++	SC
S7-SLF9 #30	S7S9	S7S7 x #12	+++	-	SI
S7-SLF9 #31	S7S11	S7S7 x #13	++	-	SI
S7-SLF9 #32	S7S17	S7S7 x #18	+++	-	SI

S7-SLF9 #33	<i>S7S19</i>	<i>S7S7</i> x #18	+++	+++	SC
S7-SLF9 #34	<i>S7S9</i>	<i>S7S7</i> x #19	+++	-	SI
S7-SLF9 #35	<i>S7S11</i>	<i>S7S7</i> x #21	++	-	SI
S7-SLF9 #36	<i>S7S17</i>	<i>S7S7</i> x #21	+++	-	SI

S11-SLF9 #1	<i>S5S17</i>	T0	+++	-	SI
S11-SLF9 #2	<i>S5S17</i>	T0	+++	-	SI
S11-SLF9 #3	<i>S5S11</i>	T0	+++	-	SI
S11-SLF9 #4	<i>S5S9</i>	<i>S9S11</i> x #1	+++	-	SI
S11-SLF9 #5	<i>S5S11</i>	<i>S9S11</i> x #1	+++	-	SI
S11-SLF9 #6	<i>S9S17</i>	<i>S9S11</i> x #1	+++	-	SI
S11-SLF9 #7	<i>S11S17</i>	<i>S9S11</i> x #1	++	-	SI
S11-SLF9 #8	<i>S5S7</i>	<i>S7S19</i> x #1	+++	-	SI
S11-SLF9 #9	<i>S5S19</i>	<i>S7S19</i> x #1	+++	+++	SC
S11-SLF9 #10	<i>S17S19</i>	<i>S7S19</i> x #1	+++	+++	SC
S11-SLF9 #11	<i>S5S11</i>	<i>S9S11</i> x #2	+++	-	SI
S11-SLF9 #12	<i>S11S17</i>	<i>S9S11</i> x #2	+++	-	SI
S11-SLF9 #13	<i>S5S7</i>	<i>S7S19</i> x #2	+++	-	SI
S11-SLF9 #14	<i>S5S19</i>	<i>S7S19</i> x #2	+	+++	SC
S11-SLF9 #15	<i>S5S19</i>	<i>S7S19</i> x #2	+++	+++	SC
S11-SLF9 #16	<i>S9S11</i>	<i>S5S9</i> x #3	+++	-	SI
S11-SLF9 #17	<i>S11S17</i>	<i>S5S17</i> x #3	+++	-	SI
S11-SLF9 #18	<i>S5S19</i>	<i>S7S19</i> x #3	+++	+++	SC
S11-SLF9 #19	<i>S5S19</i>	<i>S7S19</i> x #3	+++	+++	SC

^a The *S*-haplotype of each transgenic plant that is homoallelic with respect to the transgene is indicated in blue. For each self-compatible (SC) transgenic plant, the *S*-haplotype of pollen that exhibited competitive interaction with the transgene is indicated in red (see also Supplementary Table 6).

^b “+++” indicates the transcript of a transgene yielding a strong band after 26 cycles of amplification; “++” indicates the transcript of a transgene yielding a strong band after 28 cycles of amplification; “+” indicates the transcript of a transgene yielding a weak or moderate-intensity band after 28 cycles of amplification.

^c SC and SI phenotypes were judged by the number of the obtained seeds per pod after self-pollination. “+++” indicates more than 100 seeds set per pod; “++” indicates 51–100 seeds set per pod; “+” indicates 10–50 seeds set per pod; and “-” indicates fewer than 10 seeds set per pod. In this work, all self-pollination tests resulted in complete SC phenotype of “+++” or complete SI phenotype of “-”. All results were consistently reproduced from at least three independent self-pollinations.

^d These plants are derived from our previous work⁹.

Supplementary Table 5 Analyses of progenies of the transformants that exhibited breakdown of SI

Population		S-haplotype segregation				
Genetic cross	No. of progeny plants analyzed	Possible genotypes of progeny plants	Observed ratio ^a	Expected ratio ^b	Chi square	P-value
<i>S5S22</i> x <i>S5S22/S7-SLF1</i>	44	<i>S5S5/TG : S5S22/TG : S22S22/TG</i>	0: 21: 23	0: 1: 1 1: 2: 1	0.09 24.14	0.76 5.74E-06
<i>S11S22</i> x <i>S11S22/S7-SLF1</i>	30	<i>S11S11/TG : S11S22/TG : S22S22/TG</i>	0: 16: 14	0: 1: 1 1: 2: 1	0.13 13.20	0.72 1.36E-03
<i>S19S22</i> x <i>S19S22/S7-SLF1</i>	36	<i>S19S19/TG : S19S22/TG : S22S22/TG</i>	0: 17: 19	0: 1: 1 1: 2: 1	0.11 20.17	0.74 4.18E-05
<i>S5S7</i> x <i>S5S7/S5-SLF3</i>	52	<i>S5S5/TG : S5S7/TG : S7S7/TG</i>	0: 28: 24	0: 1: 1 1: 2: 1	0.31 22.46	0.58 1.33E-05
<i>S5S19</i> x <i>S5S19/S7-SLF9</i>	64	<i>S5S5/TG : S5S19/TG : S19S19/TG</i>	0: 35: 29	0: 1: 1 1: 2: 1	0.56 26.84	0.45 1.48E-06
<i>S7S19</i> x <i>S7S19/S7-SLF9</i>	47	<i>S7S7/TG : S7S19/TG : S19S19/TG</i>	0: 20: 27	0: 1: 1 1: 2: 1	1.04 32.06	0.31 1.09E-07
<i>S5S19</i> x <i>S5S19/S11-SLF9</i>	67	<i>S5S5/TG : S5S19/TG : S19S19/TG</i>	0: 32: 35	0: 1: 1 1: 2: 1	0.13 36.70	0.71 1.07E-08
<i>S17S19/S11-SLF9</i> self	36	<i>S17S17/TG : S17S19/TG : S19S19/TG</i>	0: 14: 22	0: 1: 1 1: 2: 1	1.78 28.67	0.18 5.96E-07

^a All progeny plants inherited the transgene (*TG*).

^b Upper ratios are values predicted by competitive interaction; lower ratios are values predicted by simple mendelian inheritance.

Supplementary Table 6 Sequence information of genes extracted from whole genome database

(a) SLF-related F-box genes from tomato

Gene_ID	Location	Motif	Query	E value
Solyc00g014960	SL2.50ch00:11,078,697..11,082,983	F-box	S7-SLF5	3.00E-25
Solyc00g014980	SL2.50ch00:11,087,048..11,093,479	F-box	S7-SLF13	9.00E-25
Solyc00g018770	SL2.50ch00:11,667,234..11,668,418	F-box + FBA1	S7-SLF5	2.00E-25
Solyc01g008040	SL2.50ch01:2,196,519..2,197,816	F-box + FBA3	S7-SLF5	1.00E-23
Solyc01g008660	SL2.50ch01:2,721,644..2,722,893	F-box	S7-SLF4	2.00E-32
Solyc01g049660	SL2.50ch01:45,830,690..45,831,775	FBA3 (ψ)	S7-SLF14	1.00E-101
Solyc01g055160	SL2.50ch01:50,895,887..50,897,056	F-box + FBA1	S7-SLF6	1.00E-43
Solyc01g056220	SL2.50ch01:52,558,634..52,559,465	F-box (ψ)	S7-SLF11	3.00E-80
Solyc01g056240	SL2.50ch01:52,616,290..52,616,775	F-box + FBA3 (ψ)	S7-SLF14	1.00E-44
Solyc01g056250	SL2.50ch01:52,683,255..52,683,869	F-box (ψ)	S7-SLF5	4.00E-56
Solyc01g056280	SL2.50ch01:52,762,364..52,763,564	F-box + FBA1	S7-FBX	1.00E-116
Solyc01_pseudo1	SL2.50ch01:52,780,047..52,781,174	F-box + FBA3 (ψ)	S7-SLF4	1.00E-158
Solyc01_pseudo2	SL2.50ch01:53,340,413..53,341,578	F-box + FBA3 (ψ)	S7-SLF1A	1.00E-136
Solyc01_pseudo3	SL2.50ch01:53,365,318..53,366,359	F-box + FBA3 (ψ)	S7-SLF4	1.00E-103
Solyc01g056660	SL2.50ch01:55,528,778..55,529,745	F-box + FBA1	S7-SLF9A	1.00E-125
Solyc01_pseudo6	SL2.50ch01:55,974,384..55,975,615	F-box + FBA1 (ψ)	S7-SLF8A	2.00E-34
Solyc01g057010	SL2.50ch01:57,933,154..57,934,326	F-box + FBA1	S7-SLF8A	1.00E-132
Solyc01g057190	SL2.50ch01:59,694,709..59,695,872	F-box + FBA1	S7-SLF13	1.00E-176
Solyc01_pseudo4	SL2.50ch01:60,535,674..60,536,864	F-box + FBA3	S7-SLF6	1.00E-153
Solyc01_pseudo5	SL2.50ch01:63,731,624..63,732,791	FBA3 (ψ)	S7-SLF5	2.00E-50
Solyc01g067010	SL2.50ch01:75,348,972..75,350,445	F-box + FBA3	S7-SLF3	1.00E-24
Solyc01g067030	SL2.50ch01:75,366,631..75,368,093	F-box	S7-SLF8A	1.00E-23
Solyc02g033040	SL2.50ch02:29,703,075..29,704,514	F-box + FBA1	S7-SLF13	1.00E-24
Solyc02g070600	SL2.50ch02:40,317,008..40,319,433	F-box + FBA3	S7-SLF4	6.00E-21
Solyc02g070620	SL2.50ch02:40,323,600..40,325,396	F-box + FBA3	S7-SLF8A	1.00E-21
Solyc03g046490	SL2.50ch03:12,935,141..12,936,370	F-box + FBA3	S7-SLF2	3.00E-21
Solyc04g063440	SL2.50ch04:55,565,206..55,566,423	F-box + FBA3	S7-SLF2	1.00E-31
Solyc04g064440	SL2.50ch04:55,581,853..55,583,076	F-box + FBA1	S7-SLF1A	2.00E-36
Solyc04g080320	SL2.50ch04:64,549,641..64,550,714	F-box + FBA1	S7-SLF8A	7.00E-23
Solyc05g005110	SL2.50ch05:115,571..116,788	F-box + FBA1	S7-SLF13	2.00E-28
Solyc05g007530	SL2.50ch05:2,085,583..2,086,701	F-box + FBA1	S7-SLF2	8.00E-23
Solyc06g059810	SL2.50ch06:37,714,157..37,717,327	F-box + FBA1	S7-SLF2	1.00E-21
Solyc07g044920	SL2.50ch07:57,969,707..57,970,804	F-box + FBA3	S7-SLF8A	2.00E-45
Solyc07g047760	SL2.50ch07:58,956,909..58,958,366	F-box + FBA3	S7-SLF14	9.00E-29
Solyc07g052580	SL2.50ch07:61,061,992..61,063,899	F-box + FBA1	S7-SLF8A	6.00E-20
Solyc07g055020	SL2.50ch07:63,166,956..63,175,330	F-box + FBA1	S7-SLF3	2.00E-24
Solyc09g072910	SL2.50ch09:65,599,205..65,600,644	F-box + FBA1	S7-SLF3	1.00E-21
Solyc09g072930	SL2.50ch09:65,606,864..65,608,024	F-box + FBA1	S7-SLF2	3.00E-27
Solyc09g083230	SL2.50ch09:68,890,249..68,891,394	F-box	S7-SLF13	6.00E-28
Solyc09g091220	SL2.50ch09:70,525,090..70,526,286	F-box	S7-SLF5	2.00E-23
Solyc09g091690	SL2.50ch09:70,938,293..70,939,459	F-box	S7-SLF8A	2.00E-31
Solyc09g091710	SL2.50ch09:70,946,048..70,947,256	F-box	S7-SLF6	3.00E-28
Solyc10g008450	SL2.50ch10:2,574,498..2,575,547	F-box + FBA1	S7-SLF8A	2.00E-20
Solyc10g008580	SL2.50ch10:2,686,335..2,687,492	F-box + FBA3	S7-SLF4	2.00E-34
Solyc10g052710	SL2.50ch10:53,260,744..53,261,892	F-box + FBA1	S7-SLF3	2.00E-21
Solyc11g012540	SL2.50ch11:5,353,456..5,355,147	F-box	S7-SLF8A	2.00E-21
Solyc11g012550	SL2.50ch11:5,357,146..5,365,080	F-box	S7-SLF8A	2.00E-23
Solyc12g006130	SL2.50ch12:690,070..691,290	F-box + FBA1	S7-SLF13	3.00E-25
Solyc12g099550	SL2.50ch12:66,661,677..66,663,236	F-box + FBA1	S7-SLF17	7.00E-27

(b) S-RNase-related ribonuclease genes from tomato

Gene_ID	Location	Motif	Query	E value
Solyc01g055200	SL2.50ch01:51,700,660..51,701,422	signal + RNase T2	S7-RNase	2.00E-59
Solyc04g005640	SL2.50ch04:417,342..420,221	signal + RNase T2	S7-RNase	1.00E-19
Solyc05g007940	SL2.50ch05:2,366,284..2,369,405	signal + RNase T2	S7-RNase	2.00E-21
Solyc05g007950	SL2.50ch05:2,373,133..2,374,737	signal + RNase T2	S7-RNase	4.00E-20
Solyc06g082890	SL2.50ch06:48,522,779..48,529,899	signal + RNase T2	S7-RNase	3.00E-14
Solyc07g006570	SL2.50ch07:1,383,208..1,384,976	signal + RNase T2	S7-RNase	3.00E-13
Solyc09g020110	SL2.50ch09:18,416,997..18,428,931	signal + RNase T2	S7-RNase	9.00E-14

Supplementary Table 6 (Continued)

(c) *SLF*-related F-box genes from potato

Gene_ID	Peptide_ID	Location	Motif	Query	E value
PGSC0003DMG400017998	PGSC0003DMP400031381	ST4.03ch00:25,449,469..25,453,906	F-box	S7-SLF5	1.00E-26
PGSC0003DMG400003427	PGSC0003DMP400006117	ST4.03ch00:29,442,018..29,443,907	F-box	S7-SLF3	5.00E-23
PGSC0003DMG400003428	PGSC0003DMP400006118	ST4.03ch00:29,447,942..29,449,474	F-box	S7-SLF3	9.00E-23
PGSC0003DMG400012104	PGSC0003DMP400021387	ST4.03ch00:38,789,791..38,791,074	F-box + FBA1	S7-SLF16	1.00E-106
PGSC0003DMG400016368	PGSC0003DMP400028605	ST4.03ch01:3,708,814..3,710,345	F-box	S7-SLF5	1.00E-22
PGSC0003DMG400008762	PGSC0003DMP400015399	ST4.03ch01:26,225,626..26,226,959	F-box + FBA1	S7-SLF8A	1.00E-132
PGSC0003DMG400009445	PGSC0003DMP400016698	ST4.03ch01:26,409,150..26,410,277	F-box + FBA1	S7-SLF4	1.00E-121
PGSC0003DMG400009446	PGSC0003DMP400016699	ST4.03ch01:26,846,171..26,847,775	F-box + FBA1	S7-SLF2	1.00E-111
PGSC0003DMG400026599	PGSC0003DMP400046220	ST4.03ch01:28,485,606..28,486,790	F-box	S7-SLF11	1.00E-125
PGSC0003DMG400006514	PGSC0003DMP400011540	ST4.03ch01:28,825,314..28,825,912	F-box + FBA1	S7-SLF14	1.00E-46
PGSC0003DMG400006512	PGSC0003DMP400011538	ST4.03ch01:28,898,538..28,899,707	F-box + FBA3	S7-SLF1A	1.00E-158
PGSC0003DMG400006511	PGSC0003DMP400011537	ST4.03ch01:28,953,578..28,954,744	F-box + FBA1	S7-SLF2	1.00E-124
PGSC0003DMG400010961	PGSC0003DMP400019375	ST4.03ch01:29,555,279..29,556,511	F-box + FBA1	S7-SLF8A	3.00E-95
PGSC0003DMG400029040	PGSC0003DMP400050586	ST4.03ch01:30,452,407..30,453,378	F-box + FBA1	S7-SLF9	1.00E-122
PGSC0003DMG400020000	PGSC0003DMP400034705	ST4.03ch01:32,070,813..32,071,627	F-box + FBA1 (ψ)	S7-SLF14	1.00E-100
PGSC0003DMG400045078	PGSC0003DMP400067182	ST4.03ch01:32,798,728..32,799,843	F-box + FBA1	S7-SLF4	7.00E-21
PGSC0003DMG400038641	PGSC0003DMP400060745	ST4.03ch01:32,916,913..32,918,037	F-box + FBA3	S7-SLF4	1.00E-125
PGSC0003DMG400014981	PGSC0003DMP400026348	ST4.03ch01:33,421,528..33,422,865	F-box + FBA3	S7-SLF6	1.00E-136
PGSC0003DMG400016927	PGSC0003DMP400029579	ST4.03ch01:40,763,687..40,764,427	F-box + FBA1	S7-SLF8A	2.00E-88
PGSC0003DMG400022664	PGSC0003DMP400039281	ST4.03ch01:58,222,546..58,224,447	F-box + FBA1	S7-SLF5	1.00E-23
PGSC0003DMG400022858	PGSC0003DMP400039629	ST4.03ch01:60,036,805..60,038,756	F-box + FBA3	S7-SLF9	4.00E-27
PGSC0003DMG400006277	PGSC0003DMP400011123	ST4.03ch01:60,245,380..60,247,024	F-box + FBA3	S7-SLF2	1.00E-22
PGSC0003DMG400006293	PGSC0003DMP400011158	ST4.03ch01:60,422,085..60,423,307	F-box + FBA1	S7-SLF3	3.00E-21
PGSC0003DMG400021478	PGSC0003DMP400037241	ST4.03ch02:17,440,094..17,444,430	F-box + FBA1	S7-SLF5	1.00E-22
PGSC0003DMG400001294	PGSC0003DMP400002321	ST4.03ch02:46,711,089..46,712,488	F-box	S7-SLF13	7.00E-25
PGSC0003DMG400039355	PGSC0003DMP400061459	ST4.03ch04:57,788,313..57,789,497	F-box	S7-SLF2	5.00E-29
PGSC0003DMG400024833	PGSC0003DMP400043043	ST4.03ch04:57,810,515..57,811,675	F-box + FBA3	S7-SLF2	3.00E-29
PGSC0003DMG400024829	PGSC0003DMP400043032	ST4.03ch04:57,860,547..57,862,051	F-box	S7-SLF2	1.00E-23
PGSC0003DMG400024825	PGSC0003DMP400043028	ST4.03ch04:57,890,755..57,892,929	F-box	S7-SLF2	1.00E-25
PGSC0003DMG401023052	PGSC0003DMP400039944	ST4.03ch05:5,462,979..5,464,210	F-box	S7-SLF13	8.00E-23
PGSC0003DMG400013491	PGSC0003DMP400023858	ST4.03ch05:9,172,804..9,178,547	F-box + FBA1	S7-SLF13	8.00E-28
PGSC0003DMG400013520	PGSC0003DMP400023917	ST4.03ch05:9,273,070..9,278,152	F-box + FBA1	S7-SLF9	4.00E-29
PGSC0003DMG400013111	PGSC0003DMP400023213	ST4.03ch05:49,049,885..49,051,148	F-box	S7-SLF9	3.00E-25
PGSC0003DMG400030753	PGSC0003DMP400053555	ST4.03ch06:41,330,354..41,331,682	F-box + FBA3	S7-SLF13	5.00E-22
PGSC0003DMG400030752	PGSC0003DMP400053554	ST4.03ch06:41,337,764..41,338,969	F-box	S7-SLF2	2.00E-21
PGSC0003DMG400013031	PGSC0003DMP400023092	ST4.03ch06:41,352,637..41,353,830	F-box + FBA1	S7-SLF2	7.00E-21
PGSC0003DMG400040257	PGSC0003DMP400062361	ST4.03ch07:3,828,392..3,829,411	F-box	S7-SLF5	1.00E-20
PGSC0003DMG400021156	PGSC0003DMP400036681	ST4.03ch07:42,476,500..42,485,182	F-box + FBA1	S7-SLF2	9.00E-33
PGSC0003DMG400020424	PGSC0003DMP400035471	ST4.03ch07:50,094,896..50,104,856	F-box	S7-SLF2	2.00E-28
PGSC0003DMG402020445	PGSC0003DMP400035515	ST4.03ch07:50,126,875..50,132,629	F-box + FBA1	S7-SLF5	8.00E-24
PGSC0003DMG400043993	PGSC0003DMP400066097	ST4.03ch07:54,136,309..54,137,370	F-box + FBA3	S7-SLF3	5.00E-23
PGSC0003DMG400022273	PGSC0003DMP400038588	ST4.03ch07:55,683,647..55,690,807	F-box	S7-SLF1A	2.00E-24
PGSC0003DMG400018496	PGSC0003DMP400032228	ST4.03ch08:3,886,902..3,888,342	F-box + FBA1	S7-SLF16	5.00E-28
PGSC0003DMG400012267	PGSC0003DMP400021719	ST4.03ch08:56,165,635..56,166,828	F-box	S7-SLF5	3.00E-25
PGSC0003DMG400002699	PGSC0003DMP400004838	ST4.03ch09:3,209,305..3,210,537	F-box	S7-SLF2	4.00E-23
PGSC0003DMG400017667	PGSC0003DMP400030874	ST4.03ch09:43,584,212..43,586,601	F-box	S7-SLF6	2.00E-22
PGSC0003DMG400003796	PGSC0003DMP400006766	ST4.03ch09:50,930,115..50,932,391	F-box + FBA1	S7-SLF2	7.00E-23
PGSC0003DMG400003797	PGSC0003DMP400006767	ST4.03ch09:50,943,971..50,946,723	F-box	S7-SLF2	5.00E-26
PGSC0003DMG400003831	PGSC0003DMP400006820	ST4.03ch09:50,974,797..50,975,936	F-box + FBA1	S7-SLF2	8.00E-26
PGSC0003DMG401006445	PGSC0003DMP400011435	ST4.03ch09:54,462,904..54,469,210	F-box	S7-SLF3	5.00E-23
PGSC0003DMG400006107	PGSC0003DMP400010841	ST4.03ch09:55,170,298..55,171,874	F-box	S7-SLF9	2.00E-23
PGSC0003DMG400029654	PGSC0003DMP400051661	ST4.03ch09:58,511,594..58,513,725	F-box + FBA1	S7-SLF5	1.00E-20
PGSC0003DMG400029632	PGSC0003DMP400051618	ST4.03ch09:58,571,145..58,573,019	F-box	S7-SLF2	1.00E-26
PGSC0003DMG400029655	PGSC0003DMP400051662	ST4.03ch09:58,581,556..58,582,764	F-box	S7-SLF5	9.00E-26
PGSC0003DMG400031300	PGSC0003DMP400054533	ST4.03ch09:58,849,836..58,850,966	F-box	S7-SLF2	4.00E-27
PGSC0003DMG400031292	PGSC0003DMP400054521	ST4.03ch09:58,851,437..58,854,176	F-box	S7-SLF5	7.00E-23
PGSC0003DMG400021673	PGSC0003DMP400037580	ST4.03ch10:4,120,452..4,121,979	F-box + FBA1	S7-SLF5	1.00E-30
PGSC0003DMG400021672	PGSC0003DMP400037579	ST4.03ch10:4,123,154..4,124,335	F-box	S7-SLF5	7.00E-28
PGSC0003DMG400031233	PGSC0003DMP400054430	ST4.03ch10:48,844,750..48,845,709	F-box + FBA3	S7-SLF13	7.00E-24
PGSC0003DMG400028237	PGSC0003DMP400049085	ST4.03ch10:56,055,611..56,057,397	F-box + FBA1	S7-SLF9	7.00E-23
PGSC0003DMG400046761	PGSC0003DMP400068865	ST4.03ch12:995,124..996,059	F-box + FBA1	S7-SLF2	1.00E-23
PGSC0003DMG400004294	PGSC0003DMP400007644	ST4.03ch12:6,242,643..6,244,440	F-box + FBA1	S7-SLF3	2.00E-21
PGSC0003DMG400045722	PGSC0003DMP400067826	ST4.03ch12:20,809,067..20,810,263	F-box	S7-SLF3	5.00E-22
PGSC0003DMG400040830	PGSC0003DMP400062934	ST4.03ch12:27,411,668..27,412,774	F-box	S7-SLF5	3.00E-21
PGSC0003DMG400034923	PGSC0003DMP400057027	ST4.03ch12:53,231,209..53,232,237	F-box	S7-SLF13	6.00E-22
PGSC0003DMG400004709	PGSC0003DMP400008352	ST4.03ch12:60,211,092..60,212,744	F-box + FBA1	S7-SLF5	6.00E-27

(d) *S-RNase*-related ribonuclease genes from potato

Gene_ID	Peptide_ID	Location	Motif	Query	E value
PGSC0003DMG400026738	PGSC0003DMP400046458	ST4.03ch00:3,948,530..3,949,620	signal + RNase T2	S7-RNase	1.00E-46
PGSC0003DMG400031503	PGSC0003DMP400054831	ST4.03ch00:35,480,463..35,481,718	RNase T2	S7-RNase	8.00E-11
PGSC0003DMG400002857	PGSC0003DMP400005163	ST4.03ch04:262,942..265,137	RNase T2	S7-RNase	1.00E-17
PGSC0003DMG400019997	PGSC0003DMP400034699	ST4.03ch05:2,484,873..2,485,866	signal + RNase T2	S7-RNase	8.00E-12

Gray shading highlights *S*-genes that belong to Solanaceae *SLFs* or *S-RNases* clades (see Fig. 3).

Supplementary table 7 Sequence information of SLF-related F-box genes, extracted from published data

Gene	Accession ID	Source	References	Note
Solanaceae SLFs				
<i>PiS1-SLF1</i>	AY500390	<i>Petunia inflata</i>	ref. 22	synonymous with PiSLF1
<i>PiS1-SLF2</i>	EF614191	<i>Petunia inflata</i>	ref. 56	synonymous with PiSLFLc-S1
<i>PiS1-SLF7</i>	EF614190	<i>Petunia inflata</i>	ref. 56	synonymous with PiSLFLa-S1
<i>PiS1-SLF9</i>	AY363970	<i>Petunia inflata</i>	ref. 37	synonymous with S1-A113
<i>PiS1-SLF10</i>	AY363973	<i>Petunia inflata</i>	ref. 37	synonymous with S1-A134
<i>PiS2-SLF1</i>	AY500391	<i>Petunia inflata</i>	ref. 22	synonymous with PiSLF2
<i>PiS2-SLF3</i>	EF614187	<i>Petunia inflata</i>	ref. 56	synonymous with PiSLFLd-S2
<i>PiS2-SLF4</i>	KF524351	<i>Petunia inflata</i>	ref. 23	
<i>PiS2-SLF5</i>	KF524352	<i>Petunia inflata</i>	ref. 23	
<i>PiS2-SLF6</i>	KF524353	<i>Petunia inflata</i>	ref. 23	
<i>PiS2-SLF7</i>	EF614189	<i>Petunia inflata</i>	ref. 56	synonymous with PiSLFLa-S2
<i>PiS2-SLF8</i>	EF614188	<i>Petunia inflata</i>	ref. 56	synonymous with PiSLFLb-S2
<i>PiS2-SLF9</i>	AY363971	<i>Petunia inflata</i>	ref. 37	synonymous with S2-A113
<i>PiS2-SLF10</i>	AY363974	<i>Petunia inflata</i>	ref. 37	synonymous with S2-A134
<i>PiS3-SLF1</i>	AY500392	<i>Petunia inflata</i>	ref. 22	synonymous with PiSLF3
<i>PiS3-SLF5</i>	KF524354	<i>Petunia inflata</i>	ref. 23	
<i>PiS3-SLF6</i>	KF524355	<i>Petunia inflata</i>	ref. 23	
<i>PiS3-SLF9</i>	AY363972	<i>Petunia inflata</i>	ref. 37	synonymous with S3-A113
<i>PiS3-SLF10</i>	AY363975	<i>Petunia inflata</i>	ref. 37	synonymous with S3-A134
<i>PiS5-SLF1</i>	KC590092	<i>Petunia inflata</i>	ref. 65	
<i>PiS7-SLF1</i>	KC590093	<i>Petunia inflata</i>	ref. 65	
<i>PiS11-SLF1</i>	KC590094	<i>Petunia inflata</i>	ref. 65	
<i>PiS13-SLF1</i>	KC590095	<i>Petunia inflata</i>	ref. 65	
<i>NaDD1-S1</i>	EF420251	<i>Nicotiana glauca</i>	ref. 66	
<i>NaDD2-S1</i>	EF420252	<i>Nicotiana glauca</i>	ref. 66	
<i>NaDD3-S1</i>	EF420253	<i>Nicotiana glauca</i>	ref. 66	
<i>NaDD4-S2</i>	EF420254	<i>Nicotiana glauca</i>	ref. 66	
<i>NaDD5-S2</i>	EF420255	<i>Nicotiana glauca</i>	ref. 66	
<i>NaDD6-S2</i>	EF420256	<i>Nicotiana glauca</i>	ref. 66	
<i>NaDD7-S2</i>	EF420257	<i>Nicotiana glauca</i>	ref. 66	
<i>NaDD8-S2</i>	EF420258	<i>Nicotiana glauca</i>	ref. 66	
<i>NaDD9-S2</i>	EF420259	<i>Nicotiana glauca</i>	ref. 66	
<i>NaDD10-S6</i>	EF420260	<i>Nicotiana glauca</i>	ref. 66	
Antirrhinum SLFs				
<i>AhSLF-S2</i>	AJ297974	<i>Antirrhinum hispanicum</i>	ref. 67	
<i>AhSLF-S2L</i>	AJ297975	<i>Antirrhinum hispanicum</i>	ref. 67	
<i>AhSLF-S1</i>	AJ515535	<i>Antirrhinum hispanicum</i>	ref. 68	
<i>AhSLF-S4</i>	AJ515534	<i>Antirrhinum hispanicum</i>	ref. 68	
<i>AhSLF-S5</i>	AJ515536	<i>Antirrhinum hispanicum</i>	ref. 68	
<i>AhSLF-S1E</i>	AJ515535	<i>Antirrhinum hispanicum</i>	ref. 68	
<i>AhSLF-S2C</i>	DQ462204	<i>Antirrhinum hispanicum</i>	ref. 68	
<i>AhSLF-S4A</i>	AJ515534	<i>Antirrhinum hispanicum</i>	ref. 68	
<i>AhSLF-S4D</i>	AJ515534	<i>Antirrhinum hispanicum</i>	ref. 68	
<i>AhSLF-S5A</i>	AJ515536	<i>Antirrhinum hispanicum</i>	ref. 68	
Maloideae SLFs/SFBBs				
<i>MdFBX1-S3</i>	AB539844	<i>Malus x domestica</i>	ref. 69	
<i>MdFBX2-S3</i>	AB539845	<i>Malus x domestica</i>	ref. 69	
<i>MdFBX3-S3</i>	AB539846	<i>Malus x domestica</i>	ref. 69	
<i>MdFBX5-S3</i>	AB539848	<i>Malus x domestica</i>	ref. 69	
<i>MdFBX7-S3</i>	AB539850	<i>Malus x domestica</i>	ref. 69	
<i>MdFBX8-S3</i>	AB539851	<i>Malus x domestica</i>	ref. 69	
<i>MdFBX9-S3</i>	AB539852	<i>Malus x domestica</i>	ref. 69	
<i>MdFBX11-S3</i>	AB539854	<i>Malus x domestica</i>	ref. 69	

<i>MdFBX14-S3</i>	AB539857	<i>Malus x domestica</i>	ref. 69	
<i>MdFBX18-S3</i>	AB539861	<i>Malus x domestica</i>	ref. 69	
<i>MdFBX20-S3</i>	AB539863	<i>Malus x domestica</i>	ref. 69	
<i>MdSFBB3-alpha</i>	AB270795	<i>Malus x domestica</i>	ref. 70	
<i>MdSFBB3-beta</i>	AB270796	<i>Malus x domestica</i>	ref. 70	
<i>PpS4FBX0</i>	AB308360	<i>Pyrus pyrifolia</i>	ref. 71	synonymous with PpSFBB1-S4 and PpSFBB4-d1
<i>PpSFBB2-S4</i>	AB270798	<i>Pyrus pyrifolia</i>	ref. 70	synonymous with PpSFBB4-β
<i>PpSFBB3-S2</i>	AB545982	<i>Pyrus pyrifolia</i>	ref. 72	synonymous with PpSFBB2-u5
<i>PpSFBB4-S4</i>	AB621610	<i>Pyrus pyrifolia</i>	ref. 73	
<i>PpSFBB5-S4</i>	AB545981	<i>Pyrus pyrifolia</i>	ref. 72	synonymous with PpSFBB4-u2
<i>PpSFBB6-S4</i>	AB270797	<i>Pyrus pyrifolia</i>	ref. 70	synonymous with PpSFBB4-α
<i>PpSFBB7-S4</i>	AB545981	<i>Pyrus pyrifolia</i>	ref. 72	synonymous with PpSFBB4-u3
<i>PpSFBB8-S4</i>	AB270799	<i>Pyrus pyrifolia</i>	ref. 70	synonymous with PpSFBB4-γ
<i>PpSFBB4-u1</i>	AB545981	<i>Pyrus pyrifolia</i>	ref. 72	
<i>PpSFBB4-u4</i>	AB545981	<i>Pyrus pyrifolia</i>	ref. 72	
<i>PpSFBB4-d2</i>	AB545981	<i>Pyrus pyrifolia</i>	ref. 72	
Prunus SLFLs				
<i>PmS7-SLFL1</i>	AB092624	<i>Prunus mume</i>	ref. 74	
<i>PmS7-SLFL2</i>	AB092626	<i>Prunus mume</i>	ref. 74	
<i>PmS7-SLFL3</i>	AB092627	<i>Prunus mume</i>	ref. 74	
<i>PaviSLFL1-S4</i>	AB280953	<i>Prunus avium</i>	ref. 75	
<i>PaviSLFL2-S4</i>	AB280954	<i>Prunus avium</i>	ref. 75	
<i>PaviSLFL3-S4</i>	AB280955	<i>Prunus avium</i>	ref. 75	
Prunus SLF/SFB/nonS-FBs				
<i>ParmSFB24</i>	HQ615603	<i>Prunus armeniaca</i>	ref. 76	
<i>PaviSFB4</i>	AB111521	<i>Prunus avium</i>	ref. 77	
<i>PdulSFB-a</i>	FJ362524	<i>Prunus dulcis</i>	Guo, C., Li, J., Luo, S. & He, T. Direct Submission	
<i>PmS7-SLF</i>	AB092622	<i>Prunus mume</i>	ref. 74	
<i>PsalSFB-c</i>	DQ849084	<i>Prunus salicina</i>	ref. 78	
<i>PaviFB</i>	JQ322648	<i>Prunus avium</i>	ref. 79	
<i>PmFbox1</i>	JX141276	<i>Prunus mume</i>	ref. 80	
<i>PmFbox2</i>	JX141277	<i>Prunus mume</i>	ref. 80	
(Outgroup)				
<i>PiFBP2011</i>	DQ250019	<i>Petunia hybrida</i>	ref. 60	
<i>PiFBP2411</i>	DQ250020	<i>Petunia hybrida</i>	ref. 60	

Supplementary table 8 Sequence information of S-RNase-related RNase-T2 genes used in phylogenetic analysis

Gene	Accession ID	Source	References	Note
Class-III: Solanaceae S-RNases				
<i>PaS1-RNase</i>	AF239908	<i>Petunia axillaris</i>	ref. 81	
<i>PaS13-RNase</i>	AF239909	<i>Petunia axillaris</i>	ref. 81	
<i>PaS15-RNase</i>	AF239910	<i>Petunia axillaris</i>	ref. 81	
<i>PaS17-RNase</i>	AY180050	<i>Petunia axillaris</i>	ref. 52	
<i>PaS19-RNase</i>	AY766156	<i>Petunia axillaris</i>	ref. 52	
<i>PaSc1-RNase</i>	AY180048	<i>Petunia axillaris</i>	ref. 52	self-compatible allele
<i>PaSc2-RNase</i>	AY180049	<i>Petunia axillaris</i>	ref. 52	self-compatible allele
<i>Pa-nonS-RNase</i>	AF239907	<i>Petunia axillaris</i>	ref. 81	non-S RNase
<i>PhS1-RNase</i>	U07362	<i>Petunia hybrida</i>	ref. 82	
<i>PhS3-RNase</i>	U07363	<i>Petunia hybrida</i>	ref. 82	
<i>PhS5-RNase</i>	AB016522	<i>Petunia hybrida</i>	ref. 51	synonymous with SB1, identical with Sv
<i>PhS7-RNase</i>	AB568388	<i>Petunia hybrida</i>	ref. 9	
<i>PhS9-RNase</i>	AB016523	<i>Petunia hybrida</i>	ref. 51	synonymous with SB2, identical with S3L
<i>PhS10-RNase</i>	AB933140	<i>Petunia hybrida</i>	This work	identical with Sx
<i>PhS11-RNase</i>	AB568389	<i>Petunia hybrida</i>	ref. 9	identical with PiS11
<i>PhS22-RNase</i>	AB933141	<i>Petunia hybrida</i>	This work	
<i>PhS22m-RNase</i>	AB933142	<i>Petunia hybrida</i>	This work	self-compatible allele
<i>PhSm-RNase</i>	AB933143	<i>Petunia hybrida</i>	This work	self-compatible allele
<i>PhS0m-RNase</i>	AB933144	<i>Petunia hybrida</i>	This work	identical with PhS0, self-compatible allele
<i>PiS1-RNase</i>	M67990	<i>Petunia inflata</i>	ref. 83	
<i>PiS2-RNase</i>	AF301533	<i>Petunia inflata</i>	ref. 84	
<i>PiS3-RNase</i>	M67991	<i>Petunia inflata</i>	ref. 83	
<i>PiS6-RNase</i>	AF301167	<i>Petunia inflata</i>	ref. 40	
<i>PiS7-RNase</i>	AF301168	<i>Petunia inflata</i>	ref. 40	
<i>PiS8-RNase</i>	AF301169	<i>Petunia inflata</i>	ref. 40	
<i>PiS9-RNase</i>	AF301170	<i>Petunia inflata</i>	ref. 40	
<i>PiS10-RNase</i>	AF301171	<i>Petunia inflata</i>	ref. 40	
<i>PiS12-RNase</i>	AF301173	<i>Petunia inflata</i>	ref. 40	
<i>PiS13-RNase</i>	AF301174	<i>Petunia inflata</i>	ref. 40	
<i>PiS15-RNase</i>	AF301175	<i>Petunia inflata</i>	ref. 40	
<i>PiS16-RNase</i>	AF301176	<i>Petunia inflata</i>	ref. 40	
<i>PiS17-RNase</i>	AF301177	<i>Petunia inflata</i>	ref. 40	
<i>PiS19-RNase</i>	AF301178	<i>Petunia inflata</i>	ref. 40	
<i>PiS20-RNase</i>	AF301179	<i>Petunia inflata</i>	ref. 40	
<i>PiS21-RNase</i>	AF301180	<i>Petunia inflata</i>	ref. 40	
<i>PiSk-RNase</i>	AB094600	<i>Petunia inflata</i>	ref. 85	
<i>PiRX2</i>	M93418	<i>Petunia inflata</i>	ref. 86	non-S RNase
<i>ScS11-RNase</i>	S69589	<i>Solanum chacoense</i>	ref. 87	
<i>ScS12-RNase</i>	AF176533	<i>Solanum chacoense</i>	ref. 88	
<i>ScS13-RNase</i>	L36667	<i>Solanum chacoense</i>	ref. 89	
<i>ScS14-RNase</i>	AF232304	<i>Solanum chacoense</i>	ref. 90	
<i>SchilS1-RNase</i>	AB072469	<i>Solanum chilense</i>	ref. 91	
<i>Sh_hab-1</i>	GU361144	<i>Solanum habrochaites</i>	ref. 92	
<i>Sh_hab-2</i>	GU361145	<i>Solanum habrochaites</i>	ref. 92	
<i>Sh_hab-3</i>	GU361146	<i>Solanum habrochaites</i>	ref. 92	
<i>Sh_hab-4</i>	GU361147	<i>Solanum habrochaites</i>	ref. 92	
<i>Sh_hab-5</i>	GU361148	<i>Solanum habrochaites</i>	ref. 92	
<i>Sh_hab-6</i>	GU361149	<i>Solanum habrochaites</i>	ref. 92	
<i>Sh_hgSRN1</i>	AB072478	<i>Solanum habrochaites</i>	ref. 91	synonymous with LhgSRN-1
<i>Sn_LpfSRN1</i>	AB072475	<i>Solanum neorickii</i>	ref. 91	synonymous with LpfSRN-1
<i>Spen_pen-1</i>	GU361150	<i>Solanum pennellii</i>	ref. 92	
<i>SpS6-RNase</i>	Z26583	<i>Solanum peruvianum</i>	ref. 93	
<i>SpS7-RNase</i>	Z26582	<i>Solanum peruvianum</i>	ref. 93	
<i>SpS11-RNase</i>	U28795	<i>Solanum peruvianum</i>	ref. 94	
<i>SpS12-RNase</i>	U28796	<i>Solanum peruvianum</i>	ref. 94	

<i>SpS13-RNase</i>	D17325	<i>Solanum peruvianum</i>	ref. 95	
<i>SpS15-RNase</i>	AB072457	<i>Solanum peruvianum</i>	ref. 91	
<i>SpS24-RNase</i>	AB072466	<i>Solanum peruvianum</i>	ref. 91	
<i>SpS25-RNase</i>	AB072467	<i>Solanum peruvianum</i>	ref. 91	
<i>Sp-nonS</i>	Z26581	<i>Solanum peruvianum</i>	ref. 93	non-S RNase
<i>StS2-RNase</i>	X62727	<i>Solanum tuberosum</i>	ref. 96	
PGSC0003DMG400026738		<i>Solanum tuberosum</i>	ref. 12	
Solyc01g055200		<i>Solanum lycopersicum</i>	ref. 13	self-compatible allele
<i>NaS2-RNase</i>	X03803	<i>Nicotiana glauca</i>	ref. 97	
<i>NaS3-RNase</i>	U66427	<i>Nicotiana glauca</i>	ref. 98	
<i>NaS6-RNase</i>	U08861	<i>Nicotiana glauca</i>	ref. 99	
<i>NaS7-RNase</i>	U13255	<i>Nicotiana glauca</i>	ref. 100	
<i>NaSA2-RNase</i>	U45957	<i>Nicotiana glauca</i>	ref. 101	
<i>Na_D63887</i>	D63887	<i>Nicotiana glauca</i>	Norioka, S. Direct Submission	
<i>Na_D63888</i>	D63888	<i>Nicotiana glauca</i>	Norioka, S. Direct Submission	
<i>Ns_relicRNase</i>	AJ002296	<i>Nicotiana glauca</i>	ref. 102	

Class-III: Antirrhinum S-RNases

<i>AhS1-RNase</i>	HE805271	<i>Antirrhinum hispanicum</i>	Liu, W. & Xue, Y. B. Direct Submission	
<i>AhS2-RNase</i>	X96465	<i>Antirrhinum hispanicum</i>	ref. 103	
<i>AhS3-RNase</i>	AJ315593	<i>Antirrhinum hispanicum</i>	Xue, Y. Direct Submission	
<i>AhS4-RNase</i>	X96466	<i>Antirrhinum hispanicum</i>	ref. 103	
<i>AhS5-RNase</i>	X96464	<i>Antirrhinum hispanicum</i>	ref. 103	

Class-III: Maloideae S-RNases

<i>MdS2-RNase</i>	U12199	<i>Malus x domestica</i>	ref. 104	
<i>MdS3-RNase</i>	U12200	<i>Malus x domestica</i>	ref. 104	
<i>MdS4-RNase</i>	AF327223	<i>Malus x domestica</i>	Van Nerum, I., et al. Direct Submission	
<i>MdS9-RNase</i>	U19793	<i>Malus x domestica</i>	ref. 104	
<i>MdS10-RNase</i>	AF327221	<i>Malus x domestica</i>	Van Nerum, I., et al. Direct Submission	
<i>MdS27b-RNase</i>	AF327222	<i>Malus x domestica</i>	Van Nerum, I., et al. Direct Submission	
<i>PpS1-RNase</i>	AB002139	<i>Pyrus pyrifolia</i>	ref. 105	
<i>PpS3-RNase</i>	AB002140	<i>Pyrus pyrifolia</i>	ref. 105	
<i>PpS4-RNase</i>	AB009385	<i>Pyrus pyrifolia</i>	ref. 106	
<i>PpS5-RNase</i>	AB002141	<i>Pyrus pyrifolia</i>	ref. 105	
<i>PpS6-RNase</i>	AB002142	<i>Pyrus pyrifolia</i>	ref. 105	
<i>PpS7-RNase</i>	AB002143	<i>Pyrus pyrifolia</i>	ref. 105	
<i>PpS8-RNase</i>	AB104908	<i>Pyrus pyrifolia</i>	ref. 107	
<i>PpS9-RNase</i>	AB104909	<i>Pyrus pyrifolia</i>	ref. 108	

Class-III: Prunus S-RNases

<i>ParmS1-RNase</i>	AY587561	<i>Prunus armeniaca</i>	ref. 109	
<i>ParmS24-RNase</i>	HQ615602	<i>Prunus armeniaca</i>	ref. 76	
<i>PaviS1-RNase</i>	AJ298310	<i>Prunus avium</i>	ref. 110	
<i>PaviS4-RNase</i>	AB028154	<i>Prunus avium</i>	ref. 111	
<i>PdulSa-RNase</i>	AB026836	<i>Prunus dulcis</i>	ref. 112	
<i>PdulSb-RNase</i>	AB011469	<i>Prunus dulcis</i>	ref. 113	
<i>PdulSc-RNase</i>	AB011470	<i>Prunus dulcis</i>	ref. 113	
<i>PmS1-RNase</i>	AB101438	<i>Prunus mume</i>	ref. 114	
<i>PmS7-RNase</i>	AB092644	<i>Prunus mume</i>	ref. 74	
<i>PmS14-RNase</i>	EU020121	<i>Prunus mume</i>	Zhang, S. L., et al., Direct Submission	
<i>PmS15-RNase</i>	EU020122	<i>Prunus mume</i>	Zhang, S. L., et al., Direct Submission	
<i>PmS16-RNase</i>	EU020123	<i>Prunus mume</i>	Zhang, S. L., et al., Direct Submission	
<i>PmSf-RNase</i>	AB101437	<i>Prunus mume</i>	ref. 114	
<i>PsalSc-RNase</i>	AB084102	<i>Prunus salicina</i>	ref. 115	
<i>PsalSb-RNase</i>	AB252413	<i>Prunus salicina</i>	ref. 116	

Class-III: S-clade non-S S-RNase-like, expressing in nectary

<i>RNasePhy3</i>	GQ465919	<i>Petunia hybrida</i>	ref. 62	expressing in nectary
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<i>RNasePhy4</i>	GQ465918	<i>Petunia hybrida</i>	ref. 62	expressing in nectary
PGSC0003DMG400019997		<i>Solanum tuberosum</i>	ref. 12	
PGSC0003DMG400031503		<i>Solanum tuberosum</i>	ref. 12	
Solyc07g006570		<i>Solanum lycopersicum</i>	ref. 13	
Class-I				
<i>RNasePhy1</i>	GQ465920	<i>Petunia hybrida</i>	ref. 62	
<i>RNasePhy5</i>	GQ465917	<i>Petunia hybrida</i>	ref. 62	
<i>RNaseNE</i>	U13256	<i>Nicotiana glauca</i>	ref. 117	
<i>RNaseNGR1</i>	AB112028	<i>Nicotiana glutinosa</i>	ref. 118	synonymous with RNaseNW
<i>RNaseNGR3</i>	AB032257	<i>Nicotiana glutinosa</i>	ref. 119	
<i>RNaseNK1</i>	AB034638	<i>Nicotiana tabacum</i>	ref. 120	
<i>RNaseLE</i>	X79337	<i>Solanum lycopersicum</i>	ref. 121	
<i>RNaseLX</i>	X79338	<i>Solanum lycopersicum</i>	ref. 121	
Solyc04g005640		<i>Solanum lycopersicum</i>	ref. 13	
Solyc05g007940		<i>Solanum lycopersicum</i>	ref. 13	
Solyc05g007950		<i>Solanum lycopersicum</i>	ref. 13	
PGSC0003DMG400002857		<i>Solanum tuberosum</i>	ref. 12	
<i>Pp_non-S</i>	D49529	<i>Pyrus pyrifolia</i>	ref. 122	non-S-RNase gene
<i>RNasePD1</i>	AF227522	<i>Prunus dulcis</i>	ref. 123	
<i>RNasePD2</i>	AF202030	<i>Prunus dulcis</i>	ref. 124	
Class-II				
<i>RNasePhy2</i>	GQ507487	<i>Petunia hybrida</i>	ref. 125	
<i>RNaseNGR2</i>	AB032256	<i>Nicotiana glutinosa</i>	ref. 119	
Solyc06g082890		<i>Solanum lycopersicum</i>	ref. 13	
Solyc09g020110		<i>Solanum lycopersicum</i>	ref. 13	
<i>RNaseLER</i>	AM408589	<i>Solanum lycopersicum</i>	ref. 126	
<i>AhSL28</i>	AJ489249	<i>Antirrhinum hispanicum</i>	ref. 127	S-like RNase 28
(Outgroup)				
<i>RNase T2</i>	NM_003730	<i>Homo sapiens</i>	ref. 63	
<i>RNase Rh</i>	D12476	<i>Rhizopus niveus</i>	ref. 64	

Classification is made according to refs. 61 and 62.

Supplementary Table 9 Summary of substitution rates among SLFs and S-RNases

Group		No. of genes	Ka	Ks
<i>Petunia</i> Type-1 SLFs		20	0.037	0.107
<i>Petunia</i> Type-2 SLFs		8	0.090	0.303
<i>Petunia</i> Type-3 SLFs		12	0.059	0.166
<i>Petunia</i> Type-4 SLFs		11	0.027	0.090
<i>Petunia</i> Type-5 SLFs		14	0.021	0.074
<i>Petunia</i> Type-6 SLFs		11	0.023	0.081
<i>Petunia</i> Type-7 SLFs		9	0.038	0.086
<i>Petunia</i> Type-8 SLFs		14	0.031	0.113
<i>Petunia</i> Type-9 SLFs		13	0.021	0.097
<i>Petunia</i> Type-10 SLFs		14	0.014	0.022
<i>Petunia</i> Type-11 SLFs		10	0.029	0.133
<i>Petunia</i> Type-12 SLFs		11	0.018	0.054
<i>Petunia</i> Type-13 SLFs		10	0.023	0.078
<i>Petunia</i> Type-14 SLFs		9	0.027	0.058
<i>Petunia</i> Type-15 SLFs		7	0.016	0.025
<i>Petunia</i> Type-16 SLFs		11	0.031	0.078
<i>Petunia</i> Type-17 SLFs		6	0.000	0.001
<i>Petunia</i> S-RNases		33	0.400	0.850
<i>P. hybrida</i> S5-haplotype	SLFs	17	0.349	0.752
<i>P. hybrida</i> S7-haplotype	SLFs	17	0.321	0.747
<i>P. hybrida</i> S9-haplotype	SLFs	18	0.325	0.756
<i>P. hybrida</i> S11-haplotype	SLFs	16	0.323	0.757
<i>P. axillaris</i> S17-haplotype	SLFs	18	0.334	0.762
<i>P. axillaris</i> S19-haplotype	SLFs	20	0.323	0.753
<i>Solanum</i> S-RNases		26	0.410	0.785
<i>S. tuberosum</i>	SLFs	14	0.405	0.950
<i>S. lycopersicum</i>	SLFs	13	0.480	1.019

Supplementary Table 10 Results of the recombination analysis using LDhat

Dataset	$4N_e r$	$\text{corr}(r^2, d)$	$P\text{corr}(r^2, d)$
All <i>SLFs</i>	49.721	-0.009	0.016
Type-3 <i>SLFs</i>	7.525	-0.023	0.047
Type-9 <i>SLFs</i>	2.975	-0.118	0.000
Type-9+10 <i>SLFs</i> + <i>FBXs</i> ^a	6.397	-0.094	0.000
Type-14+16+17 <i>SLFs</i>	8.776	-0.017	0.030

Recombination tests were carried out using the LDhat program¹⁹. Datasets analyzed were sequence alignments involving each type or some related types of *SLFs*. $4N_e r$, population recombination rate; $\text{corr}(r^2, d)$, correlations between r^2 and physical distance; $P\text{corr}(r^2, d)$, simulated p values based on 1,000 permutations. Only statistically significant results ($p < 0.05$) are shown. ^a"*FBXs*" in this table indicates type 9– and type 10–related, ungrouped *FBXs*: $S_{19}\text{-FBX1}$, $S_{19}\text{-FBX2}$, $S_{0m}\text{-FBX1}$, and $S_{0m}\text{-FBX2}$.

Supplementary Table 11 Summary of the pairwise detection of the gene conversion in *SLF* genes using GENECONV

Dataset		Allele involved	SimP	Begin	End	Length
only SI	all SLFs	S7-SLF2; S19-SLF2	0.0084	482	598	117
	Type-1 SLFs	n.d.				
	Type-2 SLFs	n.d.				
	Type-3 SLFs	PiS2-SLF3; S17-SLF3	0.0022	657	1233	577
		S10-SLF3; S22-SLF3	0.0315	732	1303	572
	Type-3+11+13 SLFs	PiS2-SLF3; S17-SLF3	0.0186	657	1233	577
	Type-4 SLFs	n.d.				
	Type-4+12 SLFs	n.d.				
	Type-5 SLFs	n.d.				
	Type-6 SLFs	n.d.				
	Type-8 SLFs	n.d.				
	Type-9+10 SLFs	PiS1-SLF10; S10-SLF10	0.0276	145	777	633
		PiS1-SLF10; S22-SLF10	0.0276	145	777	633
	Type-11 SLFs	n.d.				
	Type-11+13 SLFs	n.d.				
	Type-12 SLFs	n.d.				
	Type-13 SLFs	n.d.				
	Type-14+16+17 SLFs	n.d.				
SI + SC	all SLFs	S7-SLF2; S19-SLF2	0.0462	482	598	117
	Type-1 SLFs	S10-SLF1; Sm-SLF1	0.0120	1	659	659
	Type-2 SLFs	n.d.				
	Type-3 SLFs	S0m-SLF3; S9-SLF3	0.0066	507	831	325
		PiS2-SLF3; S17-SLF3	0.0037	601	1137	537
	Type-3+13 SLFs	S0m-SLF3;S9-SLF3	0.0263	510	834	325
		PiS2-SLF3;S17-SLF3	0.0143	604	1140	537
		S22-SLF13;Sm-SLF13	0.0035	1	535	535
		S19-SLF13;Sm-SLF13	0.0231	149	535	387
	Type-4 SLFs	n.d.				
	Type-5 SLFs	n.d.				
	Type-6 SLFs	n.d.				
	Type-7 SLFs	S17-SLF7 ψ ; S19-SLF7 ψ	0.0243	493	651	159
	Type-8 SLFs	S0m-SLF8; S5-SLF8	0.0340	205	461	257
	Type-9 SLFs	PiS3-SLF9; Sm-SLF9B	0.0057	484	759	276
		S22-SLF9; Sm-SLF9A	0.0057	988	1188	201
	Type-10 SLFs	n.d.				
	Type-9+10 SLFs + FBXs ^a	S0m-FBX1; S0m-FBX2	0.0046	62	152	91
		S0m-FBX1; S19-FBX2	0.0379	79	147	69
		S19-FBX1; S19-FBX2	0.0120	162	233	72
		Sm-SLF9B; PiS3-SLF10	0.0215	670	792	123
		Sm-SLF9B; S5-SLF10	0.0340	670	792	123
		S17-SLF9A; PiS3-SLF10	0.0340	693	811	119
		S17-SLF9B; S0m-FBX1	0.0439	37	171	135
		S17-SLF9B; S0m-FBX2	0.0157	62	152	91
		Sm-SLF9A; S19-FBX2	0.0247	319	404	86
		PiS3-SLF9; S19-FBX2	0.0262	319	411	93
		S11-SLF9; S19-FBX2	0.0380	322	411	90
	Type-11 SLFs	S11-SLF11; S19-SLF11	0.0450	246	374	129
		S17-SLF11; S19-SLF11	0.0004	247	401	155
	Type-12 SLFs	n.d.				
	Type-13 SLFs	S22-SLF13; Sm-SLF13	0.0001	1	532	532
		S5-SLF13; Sm-SLF13	0.0029	1	287	287
		S5-SLF13; S22-SLF13	0.0119	1	287	287
		S19-SLF13; Sm-SLF13	0.0154	149	532	384
		S0m-SLF13; S7-SLF13	0.0317	952	1085	134
	Type-14 SLFs	n.d.				
	Type-14+16+17 SLFs	n.d.				
	Type-15 SLFs	n.d.				
	Type-16 SLFs	n.d.				
	Type-17 SLFs	n.d.				

Gene conversion events were detected by the GENECONV program²⁰. Datasets analyzed were sequence alignments involving each type or some related types of *SLFs*. SimP, simulated *p* values based on 1,000 permutations. Only statistically significant results ($p < 0.05$) are shown. Begin, first nucleotide of converted region; End, last nucleotide in converted region; Length, length of converted region. ^a "FBXs" in this table indicates type 9- and type 10-related, ungrouped *FBXs*: *S19-FBX1*, *S19-FBX2*, *S0m-FBX1* and *S0m-FBX2*. "n.d." indicates that no significant results was detected.

Supplementary Table 12 Summary of the interactions between S-RNase and SLF experimentally demonstrated in *Petunia*

	SLFs	S-RNase alleles positively interacted	Number	S-RNase alleles negatively interacted	Number	Number of tested interaction	Proportion
Type 1	S5-SLF1 ^a	S9, S17	2	S5, S7, S11, S19	4	6	0.3333
	S7-SLF1 ^a	S9, S17, (S22) ^{b, d}	2	S5, S7, S11, S19	4	6	0.3333
	S9-SLF1 ^a	S17	1	S5, S7, S9, S11, S19	5	6	0.1667
	S11-SLF1 ^a	S17	1	S5, S7, S9, S11, S19	5	6	0.1667
	PIS2-SLF1 ^c	PIS1, PIS3, PIS7, PIS13	4	PIS2, PIS5, PIS11	3	7	0.5714
	PIS3-SLF1 ^c	-	0	PIS2, PIS3	2	2	0.0000
Type 2	S5-SLF2 ^a	S9, S11	2	S5, S7, S17, S19	4	6	0.3333
	S7-SLF2 ^a	S9, S11, S19	3	S5, S7, S17	3	6	0.5000
	S11-SLF2 ^a	S9	1	S5, S7, S11, S17, S19	5	6	0.1667
Type 3	S5-SLF3 ^b	S7	1	S5, S9, S11, S17, S19	5	6	0.1667
	S7-SLF3 ^b	-	0	S5, S7, S9, S11, S17, S19	6	6	0.0000
	S11-SLF3 ^a	S7	1	S5, S9, S11, S17, S19	5	6	0.1667
	S11-SLF3B ^b	-	0	S5, S7, S9, S11, S17, S19	6	6	0.0000
Type 4	PIS2-SLF4 ^c	PIS5	1	PIS2, PIS3, PIS6a, PIS7, PIS11, PIS12, PIS13	7	8	0.1250
Type 5	PIS2-SLF5 ^c	PIS12	1	PIS2, PIS3, PIS5, PIS6a, PIS7, PIS11, PIS13	7	8	0.1250
	PIS3-SLF5 ^c	-	0	PIS2, PIS3	2	2	0.0000
Type 6	PIS2-SLF6 ^c	-	0	PIS2, PIS3, PIS5, PIS6a, PIS7, PIS11, PIS12, PIS13	8	8	0.0000
	PIS3-SLF6 ^c	PIS2	1	PIS3	1	2	0.5000
Type 8	PIS2-SLF8 ^c	PIS6a	1	PIS2, PIS3, PIS5, PIS7, PIS11, PIS12, PIS13	7	8	0.1250
Type 9	S7-SLF9A ^b	S19	1	S5, S7, S9, S11, S17	5	6	0.1667
	S11-SLF9 ^b	S19	1	S5, S7, S9, S11, S17	5	6	0.1667
Type 13	S7-SLF13 ^a	-	0	S5, S7, S9, S11, S17, S19	6	6	0.0000
total			24		105	129	

^a Results cited from ref. 9. S₇-SLF13 is renamed from S₇-SLF3. (See caption of Supplementary Fig. 1.)

^b Results from this work.

^c Results cited from ref. 23. *Pi* indicates *Petunia inflata*.

^d Interaction between S₂₂-RNase and S₇-SLF1 eliminated from statistical analysis, because this results is based on the prediction and not random trial.

Supplementary Table 13 Statistical analyses of interactions between S-RNase and SLF

(a) Analysis for Bernouli simulation

	SLFs	S5	S7	S9	S11	S17	S19	PiS1	PiS2	PiS3	PiS5	PiS6	PiS7	PiS12	PiS13
Type 1	S5-SLF1	0	0	1	0	1	0								
	S7-SLF1	0	0	1	0	1	0								
	S9-SLF1	0	0	0	0	1	0								
	S11-SLF1	0	0	0	0	1	0								
	PiS2-SLF1				0			1	0	1	0		1		1
	PiS3-SLF1								0	0					
Type 2	S5-SLF2	0	0	1	1	0	0								
	S7-SLF2	0	0	1	1	0	1								
	S11-SLF2	0	0	1	0	0	0								
Type 3	S5-SLF3	0	1	0	0	0	0								
	S7-SLF3	0	0	0	0	0	0								
	S11-SLF3	0	1	0	0	0	0								
	S11-SLF3B	0	0	0	0	0	0								
Type 4	PiS2-SLF4				0				0	0	1	0	0	0	0
Type 5	PiS2-SLF5				0				0	0	0	0	0	1	0
	PiS3-SLF5								0	0					
Type 6	PiS2-SLF6				0				0	0	0	0	0	0	0
	PiS3-SLF6								1	0					
Type 8	PiS2-SLF8				0				0	0	0	1	0	0	0
Type 9	S7-SLF9	0	0	0	0	0	1								
	S11-SLF9	0	0	0	0	0	1								
Type 13	S7-SLF13	0	0	0	0	0	0								
Resulting factors for Bernouli simulation															
	Average	0.1860													
	S.D.	0.3907													
	Count	129													
	S.E.	0.0344													
	T-value	1.9787													
	C.I.	0.0681													

(b) Analysis for Monte Carlo simulation

	S5	S7	S9	S11	S17	S19	PiS1	PiS2	PiS3	PiS5	PiS6	PiS7	PiS12	PiS13	Postive	Total	Proportion
Type 1	0	0	1	0	1	0	1	0	1	0		1		1	6	12	0.5000
Type 2	0	0	1	1	0	1									3	6	0.5000
Type 3	0	1	0	0	0	0									1	6	0.1667
Type 4				0				0	0	1	0	0	0	0	1	8	0.1250
Type 5				0				0	0	0	0	0	1	0	1	8	0.1250
Type 6				0				1	0	0	0	0	0	0	1	8	0.1250
Type 8				0				0	0	0	1	0	0	0	1	8	0.1250
Type 9	0	0	0	0	0	1									1	6	0.1667
Type 13	0	0	0	0	0	0									0	6	0.0000

Negative and positive interactions between S-RNase and SLF, which were summarized in Supplementary Table 12, were expressed as 0 and 1, respectively.

Supplementary Table 14 Summary of the sequenced reads

Haprotypes	Total number of reads (bp)	Read length (bp) ^a	Mean. read length (bp)	Mean. base quality
<i>S5</i>	590,953	40 - 722	355	32.7
<i>S7</i>	634,016	40 - 1196	288	32.3
<i>S9</i>	550,736	40 - 893	352	32.3
<i>S11</i>	545,880	40 - 764	344	32.2
<i>S17</i>	550,612	40 - 1118	347	32.3
<i>S19</i>	521,109	40 - 987	343	32.1
<i>S0m</i>	101,431	40 - 1118	410	29.2

Total number of reads, read length, mean of read length, and mean of base quality score are shown.

^a The shortest and the longest lengths.

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